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(54) Title: HEMATOPOIETIN RECEPTORS HPR1 AND HPR2

(57) Abstract: This invention relates to human and murine HPR1 and human and murine HPR2 polypeptides, new members of the hematopoietin receptor polypeptide family; to methods of making such HPR1 and HPR2 polypeptides; to non-human mammals in which the endogenous genomic sequences encoding HPR1 and/or HPR2 polypeptides have been partially or completely inactivated; to methods of using HPR1 or HPR2 polypeptides to identify compounds that alter HPR1 or HPR2 polypeptide activities; and to methods of preparing medicaments for and/or treating conditions associated with hematopoietin receptor function.

# HEMATOPOIETIN RECEPTORS HPR1 AND HPR2

This application claims the benefit under 35 U.S.C. 119(e) of U.S. provisional applications Serial No. 60/238,706, filed 06 October 2000; Serial No. 60/240,476, filed 13 October 2000; and Serial No. 60/270,282, filed 20 February 2001; all of which are incorporated by reference herein.

# 10 <u>FIELD OF THE INVENTION</u>

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This invention relates to new human and murine hematopoietin receptor polypeptides HPR1 and HPR2, and to methods of making and using HPR1 and HPR2 polypeptides.

#### **BACKGROUND OF THE INVENTION**

The hematopoietin receptor polypeptides are a related group of Type I membrane protein receptors, and in some cases soluble forms of those receptors; this family of polypeptides has variously been called the cytokine receptor family or the hematopoietin receptor family. There are other families of receptors that bind cytokines or growth factors, such as the IL-1 receptor family, the TNF receptor family, and the EGF receptor family, but the hematopoietin receptor family is considered to be a distinct group or family of receptors based on certain characteristic structural features or motifs that are shared by members of this family. Some of the members of the hematopoietin receptor family are gp130, the granulocyte colony-stimulating factor receptor (GCSFR), leukemia inhibitory factor receptor (LIF-R), the alpha chains and the common beta chain of the IL-3 and IL-5 receptors, etc.; the hematopoietin receptor family contains more than 20 different polypeptides.

Common structural features of the hematopoietin receptor family of polypeptides include at least one extracellular cytokine receptor domain, which usually contains four cysteines and a WSXWS motif (where W is tryptophan, S is serine, and X indicates any amino acid), and, in most members of the family, a transmembrane and a cytoplasmic domain. The extracellular cytokine receptor domain is involved in ligand-binding activity, and the intracellular domain of a 'signaling' subfamily of hematopoietin receptors has a signal transduction function, transmitting the signal generated by ligand binding to a signal transduction pathway that results in the expression of genes involved in cell proliferation, differentiation, and/or activation. These activities of the hematopoietin receptor polypeptide family are mediated through interactions with cytokine ligands and other ligand-binding receptor molecules, with ligand binding to the cytokine receptor domain of hematopoietin receptor polypeptides and facilitating homo- or heterotypic interactions between receptor polypeptides, bringing the cytoplasmic domains of receptors into proximity with each other. Many of the cytokine ligands (such as IL-2, IL-6, or ciliary neurotrophic factor or CNTF, for example) interact with more than one type of heteromeric hematopoietin receptor complex, often with differing affinities, and "common" hematopoietin receptor polypeptides such as gp130 are involved in several different heteromeric receptor complexes that bind a variety of ligands. Because of their ligand-binding and intracellular signaling activities, hematopoietin receptor polypeptides are associated with a wide variety of conditions involving cytokine-influenced cell proliferation, differentiation, or activation. For example, interaction of the gp130 hematopoietin receptor polypeptide with its binding partners is involved in the

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normal upregulation of cardiac myocyte proliferation ("hypertrophy") in response to biomechanical stress on the heart, as lack of gp130 leads to heart failure under those conditions (Hirota *et al.*, 1999, *Cell* 97(2): 189-198). Hematopoietin receptors are also involved in the activation or stimulation of cells in response to environmental factors, for example the activation of hepatocytes in the acute-phase inflammatory response to injury (Taga and Kishimoto, 1992, *Crit Rev Immunol.* 11(5): 265-280; Neben and Turner, 1993, *Stem Cells* 11 Suppl 2: 156-162). Hematopoietin receptor family polypeptides generally are constitutively expressed in many different cell types throughout development, but the expression levels of hematopoietin receptor polypeptides may be up- or downregulated in response to stimuli, and some members of the family exhibit more restricted patterns of expression in particular tissues. Characteristics and activities of the hematopoietin receptor polypeptide family are described further in the following references, which are incorporated by reference herein: Drachman and Kaushansky, 1995, Curr Opin Hematol. 2(1): 22-28; Ihle, 1995, Nature 377(6550): 591-594; Taga and Kishimoto, 1995, Curr Opin Immunol. 7(1): 17-23; Ihle *et al.*, 1995, Annu Rev Immunol. 13: 369-398; Theze, 1994, Eur Cytokine Netw. 5(4): 353-368; Ihle et al., 1994, Trends Biochem Sci. 19(5): 222-227; Cosman, 1993, Cytokine 5(2): 95-106; and Onishi *et al.*, 1998, Int Rev Immunol. 16(5-6): 617-634.

In order to develop more effective treatments for disorders such as neurological, cardiac, hematopoietic, immunological, hepatic, and pulmonary conditions and diseases involving cell proliferation, differentiation, or activation, including neoplastic transformation or proliferation of virus-infected or cancerous cells, information is needed about previously unidentified members of the hematopoietin receptor polypeptide family.

25 <u>SUMMARY OF THE INVENTION</u>

The present invention is based upon the discovery of new human hematopoietin receptor family members, HPR1 and HPR2.

The invention provides an isolated polypeptide consisting of, consisting essentially of, or more preferably, comprising an amino acid sequence selected from the group consisting of:

(a) the amino acid sequence of SEQ ID NO:4;

(b) amino acids 56 through 77 of SEQ ID NO:1;

(c) an amino acid sequence selected from the group consisting of: amino acids 1 through 55 of SEQ ID NO:1; amino acids 5 through 40 of SEQ ID NO:2; amino acids 1 through 32 of SEQ ID NO:4; amino acids 1 through 241 of SEQ ID NO:4; amino acids 1 through 525 of SEQ ID NO:4; amino acids 20 through 32 of SEQ ID NO:4; amino acids 33 through 134 of SEQ ID NO:4; amino acids Xaa1 through Xaa2 of SEQ ID NO:4, wherein Xaa1 is selected from the group consisting of amino acids 33 through 43 of SEQ ID NO:4 and Xaa2 is selected from the group consisting of amino acids 228 through 241 of SEQ ID NO:4; amino acids 33 through 238 of SEQ ID NO:4; amino acids 33 through 525 of SEQ ID NO:4; amino acids 33 through 745 of SEQ ID NO:4; amino acids 44 through 94 of SEQ ID NO:4; amino acids 139 through 241 of SEQ ID NO:4; amino acids 242 through 326 of SEQ ID NO:4; amino acids 242 through 514 of SEQ ID NO:4; amino acids 337 through 419 of SEQ ID NO:4; amino acids 242 through 514 of SEQ ID NO:4; amino acids 337 through 419 of SEQ ID NO:4;

amino acids 433 through 514 of SEQ ID NO:4; amino acids 526 through 556 of SEQ ID NO:4; amino acids 533 through 552 of SEQ ID NO:4; amino acids 553 through 745 of SEQ ID NO:4; amino acids 557 through 745 of SEQ ID NO:4; amino acids 563 through 573 of SEQ ID NO:4; amino acids 563 through 641 of SEQ ID NO:4; amino acids 567 through 581 of SEQ ID NO:4; amino acids 588 through 639 of SEQ ID NO:4; and amino acids 631 through 641 of SEQ ID NO:4;

- (d) fragments of the amino acid sequences of any of (a)-(c) comprising at least 20 contiguous amino acids;
- (e) fragments of the amino acid sequences of any of (a)-(c) comprising at least 30 contiguous amino acids;
- (f) fragments of the amino acid sequences of any of (a)-(c) having HPR1 polypeptide activity;
- (g) fragments of the amino acid sequences of any of (a)-(c) comprising cytokine receptor domain amino acid sequences;
  - (h) an allelic variant of any of (a)-(c);

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- (i) amino acid sequences comprising at least 20 amino acids and sharing amino acid identity with the amino acid sequences of any of (a)-(h), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99%, and at least 99.5%;
- (j) an amino acid sequence of any of (a)-(i) wherein the polypeptide comprising said amino acid sequence also comprises an amino acid sequence selected from the group consisting of SEQ ID NO:10, SEQ ID NO:11, amino acids 652 though 745 of SEQ ID NO:4, a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least 20 contiguous amino acids; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least 30 contiguous amino acids; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 that is at least 25% of the length of the sequence of amino acids 652 though 745 of SEQ ID NO:4; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 that is at least 50% of the length of the sequence of amino acids 652 though 745 of SEQ ID NO:4; and a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4; and a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least one tyrosine residue;
- (k) an amino acid sequence of any of (a)-(j) wherein the polypeptide comprising said amino acid sequence does not comprise an amino acid sequence selected from the group consisting of amino acids 239 through 252 of SEQ ID NO:13; amino acids 643 through 652 of SEQ ID NO:14; and amino acids 652 through 662 of SEQ ID NO:15;
- (I) an amino acid sequence of (i)-(k), wherein a polypeptide comprising said amino acid sequence of (i)-(k) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (a)-(h); and
  - (m) an amino acid sequence of (i)-(l) having HPR1 polypeptide activity.

Preferably, such polypeptides are isolated HPR1 polypeptides or isolated polypeptides having HPR1 polypeptide activity.

Other aspects of the invention are isolated nucleic acids encoding polypeptides of the invention, with a preferred embodiment being an isolated nucleic acid consisting of, consisting essentially of, or more preferably, comprising a nucleotide sequence selected from the group consisting of:

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- (a) SEQ ID NO:3;
- (b) SEQ ID NO:5;
- (c) nucleotides 132 through 2366 of SEQ ID NO:3; and
- (d) allelic variants of (a)-(c).

An additional preferred embodiment of the invention is an isolated nucleic acid consisting of, consisting essentially of, or more preferably, comprising a nucleotide sequence selected from the group consisting of nucleotides 1 through 137 of SEQ ID NO:3, nucleotides 138 through 228 of SEQ ID NO:3, nucleotides 229 through 346 of SEQ ID NO:3, nucleotides 347 through 528 of SEQ ID NO:3, nucleotides 529 through 680 of SEQ ID NO:3, nucleotides 681 through 846 of SEQ ID NO:3, nucleotides 847 through 926 of SEQ ID NO:3, nucleotides 927 through 1143 of SEQ ID NO:3, nucleotides 1144 through 1326 of SEQ ID NO:3, nucleotides 1327 through 1428 of SEQ ID NO:3, nucleotides 1429 through 1575 of SEQ ID NO:3, nucleotides 1576 through 1716 of SEQ ID NO:3, nucleotides 1717 through 1810 of SEQ ID NO:3, nucleotides 1811 through 1892 of SEQ ID NO:3, and nucleotides 1893 through 2480 of SEQ ID NO:3.

The invention provides an isolated polypeptide consisting of, consisting essentially of, or more preferably, comprising an amino acid sequence selected from the group consisting of:

- (a) the amino acid sequence of SEQ ID NO:21;
- (b) an amino acid sequence selected from the group consisting of: amino acids 1 through 177 of SEQ ID NO:16; amino acids 216 through 245 of SEQ ID NO:16; SEQ ID NO:17; SEQ ID NO:18; and amino acids 349 through 356 of SEQ ID NO:23;

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an amino acid sequence selected from the group consisting of: amino acids 1 through 23 of SEQ ID NO:21; amino acids 1 through 124 of SEQ ID NO:21; amino acids 1 through 318 of SEQ ID NO:21; amino acids 1 through 331 of SEQ ID NO:21; amino acids 1 through 355 of SEQ ID NO:21; amino acids Xaa1 through Xaa2 of SEQ ID NO:21, wherein Xaa1 is selected from the group consisting of amino acids 24 through 30 of SEQ ID NO:21 and Xaa2 is selected from the group consisting of amino acids 115 through 124 of SEQ ID NO:21; amino acids 24 through 124 of SEQ ID NO:21; amino acids 24 through 331 of SEQ ID NO:21; amino acids 24 through 355 of SEQ ID NO:21; amino acids Xaa3 through Xaa4 of SEQ ID NO:21, wherein Xaa3 is selected from the group consisting of amino acids 125 through 133 of SEQ ID NO:21 and Xaa4 is selected from the group consisting of amino acids 309 through 331 of SEQ ID NO:21; amino acids 125 through 39 of SEQ ID NO:21; amino acids 125 through 31 of SEQ ID NO:21; amino acids 124 through 309 of SEQ ID NO:21; amino acids 224 through 320 of SEQ ID NO:21; amino acids 224 through 311 of SEQ ID NO:21; amino acids 319 through 350 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 325 through 331 of SEQ ID NO:21; amino acids 326 through 331 of SEQ ID NO:21; amino acids 326 through 331 of SEQ ID

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Xaa6 of SEQ ID NO:21, wherein Xaa5 is selected from the group consisting of amino acids 376 through 393 of SEQ ID NO:21 and Xaa6 is selected from the group consisting of amino acids 618 through 629 of SEQ ID NO:21; amino acids 376 through 629 of SEQ ID NO:21; amino acids 393 through 440 of SEQ ID NO:21; amino acids 393 through 618 of SEQ ID NO:21; and amino acids 397 through 611 of SEQ ID NO:21;

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- (d) fragments of the amino acid sequences of any of (a)-(c) comprising at least 20 contiguous amino acids;
- (e) fragments of the amino acid sequences of any of (a)-(c) comprising at least 30 contiguous amino acids;
- (f) fragments of the amino acid sequences of any of (a)-(c) having HPR2 polypeptide activity;
- (g) fragments of the amino acid sequences of any of (a)-(c) comprising cytokine receptor domain amino acid sequences;
  - (h) an allelic variant of any of (a)-(c);
- (i) amino acid sequences comprising at least 20 amino acids and sharing amino acid identity with the amino acid sequences of any of (a)-(h), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99.5%;
- an amino acid sequence of any of (a)-(i) wherein the polypeptide comprising said (i) amino acid sequence also comprises an amino acid sequence selected from the group consisting of: amino acids 1 through 177 of SEQ ID NO:16; amino acids 216 through 245 of SEQ ID NO:16; SEQ ID NO:17; SEQ ID NO:18; amino acids 349 through 356 of SEQ ID NO:23; amino acids 319 through 565 of SEO ID NO:21; amino acids Xaa5 through Xaa6 of SEO ID NO:21, wherein Xaa5 is selected from the group consisting of amino acids 376 through 393 of SEQ ID NO:21 and Xaa6 is selected from the group consisting of amino acids 618 through 629 of SEO ID NO:21; amino acids 376 through 629 of SEQ ID NO:21; amino acids 393 through 440 of SEQ ID NO:21; amino acids 393 through 618 of SEQ ID NO:21; amino acids 397 through 611 of SEQ ID NO:21; amino acids 381 though 629 of SEQ ID NO:21; a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 comprising at least 20 contiguous amino acids; a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 comprising at least 30 contiguous amino acids; a fragment of the sequence of amino acids 381 though 629 of SEO ID NO:21 that is at least 25% of the length of the sequence of amino acids 381 though 629 of SEQ ID NO:21; a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 that is at least 50% of the length of the sequence of amino acids 381 though 629 of SEQ ID NO:21; a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 comprising at least one of the following: an HPR2 Box 1 motif, an HPR2 Box 2 motif, and an HPR2 Box 3 motif; and a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 comprising at least one tyrosine residue;

(k) an amino acid sequence of any of (a)-(j) wherein the polypeptide comprising said amino acid sequence does not comprise amino acids 381 through 384 of SEQ ID NO:26;

- (l) an amino acid sequence of (i)-(k), wherein a polypeptide comprising said amino acid sequence of (i)-(k) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (a)-(h); and
- (m) an amino acid sequence of (i)-(l) having HPR2 polypeptide activity.

  Preferably, such polypeptides are isolated HPR2 polypeptides or isolated polypeptides having HPR2 polypeptide activity.

Other aspects of the invention are isolated nucleic acids encoding polypeptides of the invention, with a preferred embodiment being an isolated nucleic acid consisting of, consisting essentially of, or more preferably, comprising a nucleotide sequence selected from the group consisting of:

- (a) SEQ ID NO:19;
- (b) SEQ ID NO:20;
- (c) SEQ ID NO:22;
- (d) SEQ ID NO:24; and
- 20 (d) allelic variants of (a)-(d).

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An additional preferred embodiment of the invention is an isolated nucleic acid consisting of, consisting essentially of, or more preferably, comprising a nucleotide sequence selected from the group consisting of nucleotides 107 through 175 of SEQ ID NO:19, nucleotides 107 through 478 of SEQ ID NO:19, nucleotides 107 through 1099 of SEQ ID NO:19, nucleotides 107 through 1099 of SEQ ID NO:19, nucleotides 176 through 478 of SEQ ID NO:19, nucleotides 176 through 478 of SEQ ID NO:19, nucleotides 176 through 1099 of SEQ ID NO:19, nucleotides 176 through 1099 of SEQ ID NO:19, nucleotides 479 through 763 of SEQ ID NO:19, nucleotides 479 through 1099 of SEQ ID NO:19, nucleotides 503 through 1033 of SEQ ID NO:19, nucleotides 776 through 1066 of SEQ ID NO:19, nucleotides 776 through 1099 of SEQ ID NO:19, nucleotides 1232 through 1993 of SEQ ID NO:19, nucleotides 1283 through 1426 of SEQ ID NO:19, nucleotides 1283 through 1993 of SEQ ID NO:19, nucleotides 1283 through 1939 of SEQ ID NO:19, nucleotides 1283 through 1939 of SEQ ID NO:19, nucleotides 1285 through 1939 of SEQ ID NO:19.

The invention also provides isolated genomic nucleic acids corresponding to the nucleic acids of the invention.

Another aspect of the invention provides isolated nucleic acids, preferably having a length of at least 15 nucleotides, that hybridize under conditions of moderate stringency to the nucleic acids encoding polypeptides of the invention. In preferred embodiments of the invention, such nucleic acids encode a polypeptide having HPR1 and/or HPR2 polypeptide activity, or comprise a nucleotide sequence that shares nucleotide sequence identity with the nucleotide sequences of the nucleic acids of the invention, wherein the percent nucleotide sequence identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99.5%.

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Further provided by the invention are expression vectors and recombinant host cells comprising at least one nucleic acid of the invention, and preferred recombinant host cells wherein said nucleic acid is integrated into the host cell genome.

Also provided is a process for producing a polypeptide encoded by the nucleic acids of the invention, comprising culturing a recombinant host cell under conditions promoting expression of said polypeptide, wherein the recombinant host cell comprises at least one nucleic acid of the invention. A preferred process provided by the invention further comprises purifying said polypeptide. In another aspect of the invention, the polypeptide produced by said process is provided.

Further aspects of the invention are isolated antibodies that bind to the polypeptides of the invention, preferably monoclonal antibodies, also preferably humanized antibodies or humanized antibodies, and preferably wherein the antibody inhibits the activity of said polypeptides.

The invention additionally provides a method of designing an inhibitor of the polypeptides of the invention, the method comprising the steps of determining the three-dimensional structure of any such polypeptide, analyzing the three-dimensional structure for the likely binding sites of substrates, synthesizing a molecule that incorporates a predicted reactive site, and determining the polypeptide-inhibiting activity of the molecule.

In a further aspect of the invention, a method is provided for identifying compounds that alter HPR1 and/or HPR2 polypeptide activity comprising

- (a) mixing a test compound with a polypeptide of the invention; and
- (b) determining whether the test compound alters the HPR1 and/or HPR2 polypeptide activity of said polypeptide.

In another aspect of the invention, a method is provided identifying compounds that inhibit the binding activity of HPR1 and/or HPR2 polypeptides comprising

- (a) mixing a test compound with a polypeptide of the invention and a binding partner of said polypeptide; and
- (b) determining whether the test compound inhibits the binding activity of said polypeptide.

In preferred embodiments, the binding partner is a four alpha helix bundle cytokine; more preferably, the binding partner is selected from the group consisting of IL-6, OSM, LIF, CNTF, CLC, IL-12p35, and IL-23p19, and most preferably the binding partners are a soluble hematopoietin receptor such as EBI-3, soluble IL-6R alpha, cytokine-like factor-1 (CLF), IL-12p40, or a soluble form of HPR1 and/or HPR2 in conjunction with a four alpha helix bundle cytokine.

The invention also provides a method for increasing ligand-binding activity, comprising providing at least one compound selected from the group consisting of the polypeptides of the invention and agonists of said polypeptides; with a preferred embodiment of the method further comprising increasing said activity in a patient by administering at least one polypeptide of the invention.

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Further provided by the invention is a method for decreasing ligand-binding activity, comprising providing at least one antagonist of the polypeptides of the invention; with a preferred embodiment of the method further comprising decreasing said activity in a patient by administering at least one antagonist of the polypeptides of the invention, and with a further preferred embodiment wherein the antagonist is an antibody that inhibits the activity of any of said polypeptides.

The invention additionally provides a method for treating a cell proliferation condition comprising administering at least one compound selected from the group consisting of the polypeptides of the invention and agonists of said polypeptides; with a preferred embodiment wherein the cell proliferation condition is selected from the group consisting of pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, and osteoporosis resulting from a lack of bone-forming cells.

The invention additionally provides a method for treating a metabolic condition comprising administering at least one compound selected from the group consisting of the polypeptides of the invention and agonists of said polypeptides; with a preferred embodiment wherein the metabolic condition is obesity.

The invention additionally provides a method for treating a reproductive hormone condition comprising administering at least one compound selected from the group consisting of the polypeptides of the invention and agonists of said polypeptides; with a preferred embodiment wherein the condition is selected from the group consisting of deficient mammary development and infertility.

In other aspects of the invention, a method is provided for treating a cell proliferation condition comprising administering an antagonist of the polypeptide of the invention; with a preferred embodiment wherein the cell proliferation condition is selected from the group consisting of leukemia, tumour metastasis, and osteoporosis resulting from an excess of bone-resorbing cells.

In other aspects of the invention, a method is provided for treating a metabolic condition comprising administering an antagonist of the polypeptide of the invention; with a preferred embodiment wherein the metabolic condition is selected from the group consisting of cachexia, wasting, and AIDS-related weight loss.

In other aspects of the invention, a method is provided for treating cancer conditions stimulated by reproductive hormones comprising administering an antagonist of the polypeptide of the invention; with a preferred embodiment wherein the condition is selected from the group consisting of breast cancer and prolactinoma.

In another embodiment of the invention, methods are provided for using HPR1 and HPR2 polypeptides and antagonists thereof as adjuvants.

A further embodiment of the invention provides a use for the polypeptides of the invention in the preparation of a medicament for treating a cell proliferation condition; with a preferred embodiment wherein the cell proliferation condition is selected from the group consisting of pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, and osteoporosis.

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A further embodiment of the invention provides a use for the polypeptides of the invention in the preparation of a medicament for treating a metabolic condition; with a preferred embodiment wherein the metabolic condition is obesity.

A further embodiment of the invention provides a use for the polypeptides of the invention in the preparation of a medicament for treating a reproductive hormone condition; with a preferred embodiment wherein the condition is selected from the group consisting of deficient mammary development and infertility.

# **DETAILED DESCRIPTION OF THE INVENTION**

#### Similarities of HPR1 and HPR2 Structure to Other Hematopoietin Receptor Family Members

We have identified HPR1 and HPR2, new human hematopoietin receptor polypeptides having structural features characteristic of this polypeptide family; the amino acid sequence of an HPR1 polypeptide is provided in SEQ ID NO:4 and the amino acid sequence of three alternatively spliced forms of HPR2 polypeptide are provided in SEQ ID NOs 21, 23, and 25. We have also identified the murine homologue of human HPR1; the amino acid sequence of Mus musculus HPR1 is presented in SEQ ID NO:12. (The use of "HPR1" without a species designation refers to HPR1 polypeptides generally, for example, human and/or murine, mammalian, or vertebrate HPR1 polypeptides.) Alignments showing the sequence similarities between HPR1, HPR2, and other hematopoietin receptors are presented in Tables 1, 2, and 3 in Example 1 below.

The typical structural elements common to members of the hematopoietin receptor polypeptide family include an extracellular region comprising at least one cytokine receptor domain, and in most members of the family, a cytoplasmic region that in at least a subset of the hematopoietin receptor polypeptides comprises domains involved in intracellular signaling functions. A signal sequence is found at the N-terminus of hematopoietin receptor family polypeptides, and is followed, in N-to-C order, by an immunoglobulin (Ig)-like domain (in some members of the family), a cytokine receptor domain, three copies of a fibronectin repeat (in some members of the family), a transmembrane domain or a glycosyl-phosphatidyl inositol (GPI) linkage to the membrane (except in soluble members of the family, which in most cases are soluble splice variant forms of transmembrane or membrane-linked hematopoietin receptor polypeptides), and a cytoplasmic domain (which is not present in soluble forms). The extracellular domain of hematopoietin receptor polypeptides extends from the N terminus to the transmembrane domain of the protein, and includes the cytokine receptor domain and any Ig-like domains (approximately 100 amino acids in length) or fibronectin repeats (such as fibronectin type III repeats which are approximately 81-83 amino acids in length and are separated by spacer sequences of approximately 10 to 13 amino acids) that may be present in certain of the hematopoietin receptor polypeptides. There are key residues within the cytokine receptor domain, the two or four conserved cysteine residues and the WSXWS motif; substitutions of these residues are likely to be associated with an altered function or lack of that function for the polypeptide. The cytokine receptor domain, which is approximately 200 amino acids in length, can be subdivided into two roughly equal subdomains - an N-terminal 'conserved cysteine' domain and a more C-terminal

'WSXWS' domain - separated by a proline-rich 'linker' stretch of four amino acids that allows the two subdomains to form a ligand binding site between them (Bravo and Heath, 2000, *EMBO J.* 19(11): 2399-2411).

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The intracellular domain (also called "cytoplasmic domain") of the hematopoietin receptor polypeptides (in those family members that contain such a domain), extends from the transmembrane domain of the protein to the C terminus, and in the signaling receptor subgroup, includes regions involved in intracellular signal transduction functions. Although the amino acid sequence of the intracellular domain varies considerably between hematopoietin receptor polypeptides, there are a few regions that show some similarity between the members of the family and which have been determined to be involved in binding to members of the signal transduction cascade. "Box 1" is a stretch of 9 to 12 amino acids that begins about 9 amino acids C-terminal to the transmembrane domain, and has within it a conserved Ar-P-X-Al-P-X-P motif, where Ar is an aromatic amino acid (Trp, Phe, or Tyr) and Al is an aliphatic amino acid (Ala, Gly, Val, Leu, or Ile). About 8 amino acids C-terminal to Box 1 there is a conserved aromatic amino acid (usually Trp but also Phe or Tyr), and approximately 15 to 60 amino acids further C-terminal there is a motif of about 11 to 13 amino acids, "Box 2". While Box 1 is present in most of the hematopoietin receptor polypeptides, the Box 2 motif is present in a subset of the hematopoietin receptor family including gp130, GCSFR, LIF-R, the erythropoietin receptor (EPO-R), and several others. Mutations to residues within Box 1 or Box 2, or to the conserved aromatic residue between the Box 1 and Box 2 motifs, have inactivated the ability of the mutated receptor to stimulate cell proliferation upon the addition of ligand. A further conserved domain has been identified in the cytoplasmic domains of signaling cytokine receptors such as gp130, LIF-R, and G-CSFR: "Box 3". The Box 3 motif is about 10 to 15 amino acids located between approximately 70 and 150 amino acids C-terminal of the transmembrane domain, and has a rough match to a (P/T)VXGXGYXXQ consensus sequence. Cytoplasmic regions of these receptors containing Box 3 have been associated with a macrophage differentiation promoting activity (in the case of gp130) and a granulocyte differentiation promoting activity (in the case of G-CSFR) (Soede-Bobok and Touw, 1997, J Mol Med 75: 470-477); however, members of the LIF/IL-6 gp130-sharing family of hematopoietin receptors can also be involved in suppression of differentiation (see Ernst et al., 1999, J Biol Chem 274(14): 9729-9737). Finally, the cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation. Although hematopoietin receptors themselves do not generally have a protein kinase activity, they interact with and are phosphorylated by kinases within the JAK/STAT signal transduction pathways. Mutations in the Box 1 motif abolish the ability of certain of the signaling hematopoietin receptors to bind members of the Janus kinase (JAK) family, particularly JAK2 or JAK1 (Taner et al., 1995, J Biol Chem 270(12): 6523-6530). Hematopoietin receptor-ligand interactions also activate the ERK/MAPK pathway, most likely through the phosphorylation of tyrosine residues in the cytoplasmic domains as the tyrosines at cytoplasmic positions 118 of gp130 (amino acid 759 of SEQ ID NO:8) and 115 of LIF-R (amino acid 974 of SEQ ID NO:6) are present within SHP2 binding sites (Schiemann et al., 1997, J Biol Chem 272(26): 16631-

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16636). The cytoplasmic tyrosine residues of signaling hematopoietin receptors and the amino acids around them are also important motifs for the recruitment and phosphorylation of signal-transducing STAT polypeptides (Hirano *et al.*, 2000, *Oncogene* 19: 2548-2556).

Human HPR1 polypeptide has a signal sequence extending from approximately amino acid 20 through amino acid 32 of SEQ ID NO:4, with the mature polypeptide produced by cleavage of this signal sequence predicted to have an amino acid sequence beginning at amino acid 33 of SEQ ID NO:4. Human HPR1 has a cytokine receptor domain extending approximately from amino acid 33 through amino acid 241 of SEQ ID NO:4; three fibronectin repeats from approximately amino acid 242 of SEQ ID NO:4 to about amino acid 515 of SEQ ID NO:4; a transmembrane domain that begins approximately between amino acids 526 and 533 of SEQ ID NO:4 and extends to approximately between amino acids 552 and 556 of SEQ ID NO:4 (defining a smaller 'core' transmembrane domain from amino acid 533 to amino acid 552 of SEQ ID NO:4 and an extended transmembrane domain from amino acid 526 to amino acid 556 of SEQ ID NO:4); and a cytoplasmic domain extending from the end of the transmembrane domain (i.e. beginning roughly between amino acids 553 and 557 of SEQ ID NO:4) and extending through the carboxyl terminus of the polypeptide (amino acid 745 of SEO ID Therefore, human HPR1 polypeptide has an overall structure consistent with other hematopoietin receptor family members. The four conserved cysteine residues within the human HPR1 cytokine receptor domain are located at positions 43, 53, 81, and 94 of SEQ ID NO:4, and the human HPR1 WSXWS motif is located from amino acid 224 through amino acid 228 of SEQ ID NO:4. The human HPR1 N-terminal cytokine receptor subdomain containing four conserved cysteine residues extends approximately from amino acid 33 of SEQ ID NO:4 to amino acid 134 of SEQ ID NO:4; the proline-rich linker is amino acids 135 through 138 of SEQ ID NO:4; and the WSXWS-containing Cterminal cytokine receptor subdomain extends from amino acid 139 to about amino acid 241 of SEQ ID NO:4. In human HPR1, as in several members of the hematopoietin receptor family, the cytokine receptor domain is followed by three fibronectin type III repeats; these repeats are located within the human HPR1 amino acid sequence of SEQ ID NO:4 at the following approximate locations: amino acids 242 to 244 through 324 to 326, amino acids 336 to 337 through 419 to 422, and amino acids 430 to 433 through 514 to 515. Within its intracellular domain, human HPR1 polypeptide contains a good match to the Box 1 conserved motif from amino acid 563 through amino acid 573 of SEQ ID NO:4, a conserved downstream Trp residue (amino acid 581 of SEQ ID NO:4), and a Box 2 motif from amino acid 631 to amino acid 641 of SEQ ID NO:4. The cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation; in human HPR1, such tyrosines are located at positions 652, 683, and 721 of SEQ ID NO:4. Human HPR1 contains several instances of an Asp-containing motif within its cytoplasmic region. In the area overlapping the Box 2 location, human HPR1 has repeated amino acid sequences as shown in the following table; these sequences form a consensus sequence of DKL(N/V)(T/Al), where Al is an aliphatic residue as described above. Other signaling hematopoietin receptors such as murine HPR1 (at

5 amino acids 600 through 604 of SEQ ID NO:12) and gp130 also contain at least one similar Aspcontaining sequence in the region around and following the Box 2 location.

Repeat Sequence	Location in SEQ ID NO:4
DKLNL	amino acids 588 through 592
DSVNT	amino acids 597 through 601
DRILK	amino acids 603 through 607
DKLVI	amino acids 614 through 618
DKLVV	amino acids 619 through 623
DEART	amino acids 635 through 639

Variants, presumably splice variants, of human HPR1 are described in WO 00/75314: a 252-amino-acid form ("NR10.2"), a 652-amino-acid form ("NR10.1"), and a 662-amino-acid form ("NR10.3"). The 252-amino-acid form of HPR1 (SEQ ID NO:13) is identical to SEQ ID NO:4 through amino acid 238, and then has a divergent amino acid sequence from amino acid 239 through 252 of SEQ ID NO:13. This 252-amino-acid form of human HPR1 therefore does not contain the fibronectin type III repeats found in the full-length 745-amino-acid HPR1 of SEQ ID NO:4, or the transmembrane domain or the intracellular region of the SEQ ID NO:4 polypeptide. The 652-amino-acid form of HPR1 (SEQ ID NO:14) is identical to SEQ ID NO:4 through amino acid 642, and then has a divergent amino acid sequence from amino acid 643 through 652 of SEQ ID NO:14.; and the 662-amino-acid form of HPR1 (SEQ ID NO:15) is identical to SEQ ID NO:4 through amino acid 651, and then has a divergent amino acid sequence from amino acid 652 through 662 of SEQ ID NO:15. The 652- and 662-amino-acid forms of human HPR1 therefore do not contain the tyrosine residues at positions 652, 683, and 721 of the intracellular region of the SEQ ID NO:4 polypeptide which are potential substrates for phosphorylation by kinases, such as those of the ERK/MAPK signaling pathways.

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The Mus musculus HPR1 amino acid sequence of SEQ ID NO:12 has a signal sequence beginning approximately between amino acid 13 and amino acid 16 of SEQ ID NO:12 and extending approximately through amino acid 28 of SEQ ID NO:12, with the mature polypeptide produced by cleavage of this signal sequence predicted to have an amino acid sequence beginning at amino acid 29 of SEQ ID NO:12. Murine HPR1 has a cytokine receptor domain extending approximately from amino acid 29 through amino acid 224 of SEQ ID NO:12; three fibronectin repeats from approximately amino acid 225 of SEQ ID NO:12 to about amino acid 499 of SEQ ID NO:12; a transmembrane domain that begins approximately between amino acids 510 and 517 of SEO ID NO:12 and extends to approximately between amino acids 532 and 533 of SEQ ID NO:12 (defining a smaller 'core' transmembrane domain from amino acid 517 to amino acid 532 of SEQ ID NO:12 and an extended transmembrane domain from amino acid 510 to amino acid 533 of SEQ ID NO:12); and a cytoplasmic domain extending from the end of the transmembrane domain (i.e. beginning roughly between amino acids 533 and 534 of SEQ ID NO:12) and extending through the carboxyl terminus of the polypeptide (amino acid 726 of SEQ ID NO:12). Therefore, murine HPR1 polypeptide has an overall structure consistent with other hematopoietin receptor family members. There are two conserved cysteine residues within the murine HPR1 cytokine receptor domain located at positions 39 and 49of SEQ ID NO:12, and there are two additional cysteines in this region (although at non-conserved positions) at

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amino acids 90 and 97 of SEQ ID NO:12. The murine HPR1 WSXWS motif is located from amino acid 207 through amino acid 211 of SEQ ID NO:12. The murine HPR1 N-terminal cytokine receptor subdomain containing two conserved cysteine residues (and two additional cysteine residues) extends approximately from amino acid 29 of SEQ ID NO:12 to amino acid 124 of SEQ ID NO:12; the prolinerich linker is amino acids 125 through 128 of SEQ ID NO:12; and the WSXWS-containing C-terminal cytokine receptor subdomain extends from amino acid 129 to about amino acid 224 of SEQ ID NO:12. In murine HPR1, as in several members of the hematopoietin receptor family, the cytokine receptor domain is followed by three fibronectin type III repeats; these repeats are located within the murine HPR1 amino acid sequence of SEQ ID NO:12 at the following approximate locations: amino acids 225 to 227 through 307 to 309, amino acids 319 to 320 through 403 to 406, and amino acids 413 to 417 through 498 to 499. Within its intracellular domain, murine HPR1 polypeptide contains a good match to the Box 1 conserved motif from amino acid 547 through amino acid 557 of SEQ ID NO:12, a conserved downstream Trp residue (amino acid 565 of SEQ ID NO:12), and a Box 2 motif from amino acid 612 through amino acid 622 of SEQ ID NO:12. The cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation; in murine HPR1, such tyrosines are located at positions 633, 674, and 701 of SEQ ID NO:12.

Human HPR2 polypeptide has a signal sequence extending from approximately amino acid 11 through amino acid 23 of SEQ ID NO:21, with the mature polypeptide produced by cleavage of this signal sequence predicted to have an amino acid sequence beginning at amino acid 24 of SEQ ID NO:21. The membrane-spanning (629 amino acids) form of HPR2 has an N-terminal Ig-like domain extending approximately from amino acid 24 through amino acid 124 of SEQ ID NO:21, a cytokine receptor domain extending approximately from amino acid 125 through an amino acid from 320 to 331 of SEQ ID NO:21; a transmembrane domain that begins approximately at amino acid 356 of SEQ ID NO:21 and extends to approximately amino acid 375 of SEQ ID NO:21; and a cytoplasmic domain extending from the end of the transmembrane domain (i.e. beginning approximately at amino acid 376 of SEQ ID NO:21) and extending through the carboxyl terminus of the polypeptide (amino acid 629 of SEQ ID NO:21). Therefore, HPR2 polypeptide has an overall structure consistent with other hematopoietin receptor family members. The N-terminal Ig-like domain contains six cysteine residues at positions 30, 52, 59, 101, 105, and 115 of SEQ ID NO:21, the most conserved of which appear to be the two cysteines at positions 52 and 101; the cysteines at positions 30, 115 (and to a lesser extent, at 105) also align with cysteines at similar positions in Ig or Ig-like domains. The HPR2 Ig-like domain appears to have the greatest degree of sequence similarity with members of the LIR (leukocyte Ig-like receptor) polypeptide family, particularly LIR-3 and LIR-4. The two conserved cysteine residues within the human HPR2 cytokine receptor domain are located at amino acid positions 133 and 144 of SEQ ID NO:21, and the HPR2 version of the WSXWS motif, which has a glutamine residue at the second position of the motif rather than a serine residue, is located from amino acid 304 through amino acid 308 of SEQ ID NO:21. The HPR2 N-terminal cytokine receptor subdomain containing the two

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conserved cysteine residues extends approximately from amino acid 125 of SEQ ID NO:21 to amino acid 219 of SEQ ID NO:21; the proline-rich linker (in this case, proline- and alanine-rich) is amino acids 220 through 223 of SEO ID NO:21; and the 'WOXWS'-containing C-terminal cytokine receptor subdomain extends from amino acid 224 through an amino acid from 320 to 331 of SEQ ID NO:21. HPR2 does not contain the fibronectin type III repeats found in human and murine HPR1. Within its intracellular domain, the membrane-spanning (629 amino acids) form of HPR2 contains a good match to the Box 1 conserved motif from amino acid 393 through amino acid 403 of SEQ ID NO:21, does not contain a Trp residue between Box 1 and Box2, and has a Box 2 motif from amino acid 430 to amino acid 440 of SEQ ID NO:21. There are also two matches to the Box 3 motif in this membrane-spanning HPR2 polypeptide, at amino acids 478 through 491 and at amino acids 605 through 618 of SEQ ID NO:21. The cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation; in human HPR2, such tyrosines are located at amino acid positions 397 (within the Box 1 motif), 429 (immediately N-terminal to the Box 2 motif), 450, 463, and 476 (just N-terminal of the most N-terminal Box 3 motif), and amino acids 484 and 611 (each of these last two amino acids is within a Box 3 motif) of SEQ ID NO:21. In several respects, the membrane-spanning form of HPR2 shows similarity to the LIF-R hematopoietin receptor: both of these molecules have an Ig-like domain that is followed by a cytokine receptor domain having two (as compared to four) conserved cysteines; and both have Box 1, Box 2, and Box 3 motifs in their intracellular domains, and do not have a tryptophan residue between Box 1 and Box 2.

The HPR2-ex9 polypeptide of SEQ ID NO:23 (356 amino acids), created by alternative splicing which removes exon 9 of the HPR2 coding sequence (see Example 1 below), is identical to the HPR2 629-amino-acid form from amino acid 1 through amino acid 348, but then diverges in sequence for the eight amino acids from amino acid 349 to the C terminus at amino acid 356. The HPR2-ex9 form does not contain a transmembrane region, and is expected to be a secreted form of HPR2 containing the HPR2 extracellular Ig-like and cytokine receptor domains. The HPR2-ex8-ex9 polypeptide of SEQ ID NO:25 (565 amino acids), created by alternative splicing which removes exons 8 and 9 of the HPR2 coding sequence (see Example 1 below), is identical to the HPR2 629-amino-acid form from amino acid 1 through amino acid 318, is missing the next 64 amino acids which include the transmembrane domain, but then shows identity between amino acid 319 through amino acid 565 of SEQ ID NO:25 and the C-terminal region of the 629-amino-acid form of HPR2. The HPR2-ex8-ex9 form is also expected to be a secreted form of HPR2 containing not only the HPR2 extracellular Ig-like and cytokine receptor domains, but also the C-terminal portion of the HPR2 protein which includes the Box 1, Box 2, and Box 3 motifs. A variant, presumably a splice variant, of human HPR2 is described in WO 00/73451: a 384-amino-acid form ("DCRS2"). This 384-amino-acid form of HPR2 (SEQ ID NO:26) is identical to SEQ ID NO:21 through amino acid 380, and then has a divergent amino acid sequence from amino acid 381 through 384 of SEQ ID NO:26. This 384-amino-acid form of human HPR2 therefore does not contain the intracellular region of the SEQ ID NO:21 HPR2 polypeptide,

which contains the Box1, 2, and 3 motifs and intracellular tyrosine residues that are involved in the signaling (or signal transduction) function of the SEQ ID NO:21 HPR2 polypeptide.

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The Mus musculus HPR2 amino acid sequence of SEQ ID NO:27 has a signal sequence beginning approximately between amino acid 8 and amino acid 11 and extending through amino acid 23 of SEQ ID NO:27, with the mature polypeptide produced by cleavage of this signal sequence predicted to have an amino acid sequence beginning at amino acid 24 of SEQ ID NO:27. Mus musculus HPR2, like the membrane-spanning form of human HPR2, has an N-terminal Ig-like domain extending approximately from amino acid 24 through amino acid 124 of SEQ ID NO:27, a cytokine receptor domain extending approximately from amino acid 125 through an amino acid from 341 to 350 of SEQ ID NO:27; a transmembrane domain that begins approximately between amino acid 373 and amino acid 380 of SEQ ID NO:27 and extends through approximately between amino acid 394 and amino acid 395 of SEQ ID NO:27 (defining a smaller 'core' transmembrane domain from amino acid 380 to amino acid 394 of SEQ ID NO:27 and an extended transmembrane domain from amino acid 373 to amino acid 395 of SEQ ID NO:27); and a cytoplasmic domain extending from the end of the transmembrane domain (i.e. beginning approximately at amino acid 395 or at amino acid 396 of SEO ID NO:27) and extending through the carboxyl terminus of the polypeptide (amino acid 644 of SEQ ID Therefore, murine HPR2 polypeptide has an overall structure consistent with other hematopoietin receptor family members. The N-terminal Ig-like domain contains six cysteine residues at positions 30, 52, 59, 101, 105, and 115 of SEQ ID NO:27, the most conserved of which appear to be the two cysteines at positions 52 and 101; the cysteines at positions 30, 115 (and to a lesser extent, at 105) also align with cysteines at similar positions in Ig or Ig-like domains. As with human HPR2, the murine HPR2 Ig-like domain appears to have the greatest degree of sequence similarity with members of the LIR (leukocyte Ig-like receptor) polypeptide family. The two conserved cysteine residues within the human HPR2 cytokine receptor domain are located at amino acid positions 133 and 144 of SEQ ID NO:27, and the murine HPR2 version of the "WSXWS" motif, which like human HPR2 has a glutamine residue at the second position of the motif rather than a serine residue, is located from amino acid 324 through amino acid 328 of SEQ ID NO:27. The murine HPR2 polypeptide contains an insert of 20 amino acids relative to the human HPR2 polypeptide; this insert region extends from amino acid 297 through amino acid 316 of SEQ ID NO:27, and is a perfect repeat of amino acids 317 through 336 of SEQ ID NO:27. Therefore, in the SEQ ID NO:27 form of murine HPR2, there is a second WOXWS motif at amino acids 304 through 308 of SEQ ID NO:27. The murine HPR2 N-terminal cytokine receptor subdomain containing the two conserved cysteine residues extends approximately from amino acid 125 of SEQ ID NO:27 to amino acid 219 of SEQ ID NO:27; the proline-rich linker (in this case, proline- and alanine-rich) is amino acids 220 through 223 of SEQ ID NO:27; and the C-terminal cytokine receptor subdomain containing the two repeats of the WQXWS motif extends from amino acid 224 through an amino acid from 340 to 350 of SEQ ID NO:27. Murine HPR2 does not contain the fibronectin type III repeats found in human and murine HPR1. Within its intracellular domain, this membrane-spanning form of murine HPR2 contains a good match to the Box 1 conserved motif from

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amino acid 412 through amino acid 422 of SEQ ID NO:27, does not contain a Trp residue between Box 1 and Box2, and has a Box 2 motif from amino acid 449 to amino acid 459 of SEQ ID NO:27. There are also two matches to the Box 3 motif in this murine membrane-spanning HPR2 polypeptide, at amino acids 498 through 511 and at amino acids 620 through 633 of SEQ ID NO:27. The cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation; in murine HPR2, such tyrosines are located at amino acid positions 416 (within the Box 1 motif), 448 (immediately N-terminal to the Box 2 motif), 469, and 496 (just N-terminal of the most N-terminal Box 3 motif), and amino acids 504 and 626 (each of these last two amino acids is within a Box 3 motif) of SEQ ID NO:27. There is an additional intracellular tyrosine located at position 542 of SEQ ID NO:27. As with the membrane-spanning form of human HPR2, murine HPR2 shows similarity to the LIF-R hematopoietin receptor.

Each of the HPR1 and the HPR2 groups of related polypeptides therefore contains a distinct subset of the several features characteristic of at least some members of the hematopoietin receptor family. The skilled artisan will recognize that the boundaries of the regions of the HPR1 and HPR2 polypeptides described above are approximate and that the precise boundaries of such domains, as for example the boundaries of the transmembrane region (which can be predicted by using computer programs available for that purpose), can also differ from member to member within the hematopoietin receptor polypeptide family.

The hematopoietin receptor polypeptide family is highly to moderately conserved between species, with the family members within a particular species exhibiting some sequence conservation, particularly with respect to the conserved domains and residues described above. Subfamilies of the hematopoietin receptor polypeptide family can be defined on the basis of structure, for example the Iglike domain containing members, or the fibronectin repeat containing members. It is also possible to group hematopoietin receptor polypeptides according to the length of the cytoplasmic domain, with those receptors having a longer cytoplasmic domain being more likely to be signaling receptors. Subgroups of the hematopoietin receptor family can also be defined on the basis of a shared common signaling receptor present in several different combinations of heteromeric receptors. For example, the gp130 signaling receptor is found in separate complexes with LIF-R, IL-6R alpha or a soluble form of IL-6R alpha, and CNTFR alpha; monomeric forms or multimeric combinations of these receptor components bind to IL-6, OSM, LIF, and/or CNTF; thus a "gp130-sharing group" subfamily would include these hematopoietin receptor polypeptides and be associated with this group of cytokines. Another group of hematopoietin receptors are those which associate with a ligand comprising at least two soluble polypeptides. For example, the IL-12 receptor associates with the combination of the p40 polypeptide, similar in structure to soluble forms of hematopoietin receptors such as soluble IL-6R alpha, and the four alpha helix bundle p35 polypeptide. The IL-12 p40 subunit can also associate with another four alpha helix bundle cytokine called p19; when p40 binds p19 the resulting combination has been named "IL-23" and has been shown to bind to the IL-12R beta 1 receptor subunit, but not the signaling IL-12R beta 2 receptor subunit (Oppmann et al., 2000, Immunity 13: 715-725). Thus the

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p40-p19 complex is likely to bind a different IL-12RB2-like signaling receptor subunit, such as HPR2, HPR1, GCSFR, or gp130. As another example, CNTFR alpha, gp130, and LIFR can each associate with a combination of the soluble receptor cytokine-like factor-1 (CLF-1) and cardiotrophin-like cytokine (CLC), with CLF-1 and CLC analogous to p40 and p35, respectively (Elson et al., 2000, Nat Neurosci 3(9): 867-872). The cytokine receptor domains of HPR1 and HPR2 are similar in sequence to those of gp130, IL-6R beta, IL-12RB2, GCSFR, LIFR, leptin receptor, prolactin receptor, and other members of the hematopoietin receptor family, with HPR1 showing the greatest degree of similarity to gp130 and IL-6R beta, and HPR2 showing the greatest degree of similarity to gp130 and IL-12RB2. Because HPR1 and HPR2 each have a substantial cytoplasmic domain and are most similar in sequence to gp130, HPR1 and HPR2 are likely to be new signaling members of the "gp130-sharing" subfamily of hematopoietin receptors; however, HPR2 may also share attributes of the IL-12RB2 receptor subunit, such as involvement in modulation of the balance between Th1 and Th2 immune responses. Expression of HPR1 and HPR2 has been detected by PCR amplification from tissue-specific cDNA libraries in several cell types including COS-1 cells, 293MSR cells, the B cell lines CB23 and MP-1, the B cell lymphoma lines Daudi, and Raji, the T cell leukemia line HSB2, and the promonocytic leukemia line U937. HPR2 mRNA expression appears to be more prevalent than HPR1 expression in the B cell derived lines, while HPR1 mRNA expression appears to be more prevalent than HPR2 expression in the T cell derived and monocyte lines. EBI-3 is a p40-like soluble hematopoietin receptor polypeptide; FACS analysis has shown that EBI-3-Fc fusion polypeptides bind to cells expressing HPR1 and HPR2 such as COS-1 cells, 293MSR cells, and CB23 and MP-1 cells, indicating that EBI-3 is a potential binding partner of HPR1 and HPR2, most likely in conjunction with a four alpha helix bundle cytokine such as IL-6, OSM, LIF, CNTF, CLC, IL-12p35, or IL-23p19.

#### Biological Activities and Functions of HPR1 and HPR2 Polypeptides

PCR amplification from tissue-specific cDNA libraries was performed to detect HPR1 or HPR2 cDNA sequences. The results of these experiments show that HPR1 transcripts are expressed in a wide variety of fetal and adult human cells, including testis, lung, placenta, pancreas, prostate, peripheral blood cells, thymus, stomach, and skin cells; as well as in various cell lines including U937 cells, the leukemia cell line HSB2, LX-1/GI-117 lung carcinoma cells, GI-112 colon adenocarcinoma cells, the B cell lines MP-1 and CB23, COS-1 cells, and 293MSR cells. HPR2 transcripts are present in a similarly diverse group of adult and fetal human cell types, including placenta, lung, kidney, pancreas, prostate, testis, colon, LX-1/GI-117 lung carcinoma cells, tonsil/CX-1 cells, lymph node, GI-112 colon adenocarcinoma cells, heart, brain, spleen, thymus, ovary, small intestine, fetal brain, fetal lung/heart, fetal spleen, fetal thymus, esophagus, stomach, and skin; and in various cell lines such as the B cell lines MP-1 and CB23, Daudi cells, Raji cells, HSB2 cells, COS-1 cells, and 293MSR cells.

Typical biological activities or functions associated with HPR1 and HPR2 polypeptides are ligand-binding activity, intracellular signaling activity, cell proliferation stimulatory activity, cell proliferation inhibitory activity, cell differentiation stimulatory activity, and cell differentiation

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inhibitory activity. HPR1 and HPR2 polypeptides having ligand-binding activity bind to cytokine or growth factor ligand molecules of the four alpha helix bundle family of cytokines, and in particular are likely to bind cytokines such as IL-6, OSM, LIF, CNTF, CLC, IL-12p35, and IL-23p19, and/or soluble hematopoietin receptors such as EBI-3, soluble IL-6R alpha, cytokine-like factor-1 (CLF), IL-12p40, or a soluble form of HPR1 and/or HPR2. This ligand-binding activity is associated with the extracellular cytokine receptor domain of HPR1 polypeptides. Thus, for uses requiring ligand-binding activity, preferred HPR1 and HPR2 polypeptides include those having at least one cytokine receptor domain and exhibiting ligand-binding activity. Preferred HPR1 and HPR2 polypeptides further include oligomers or fusion polypeptides comprising at least one cytokine receptor portion of one or more HPR1 and/or HPR2 polypeptides, and fragments of any of these polypeptides that have ligand-binding activity. The ligand-binding activity of HPR1 and HPR2 polypeptides may be determined, for example, by any standard assay to measure binding of labeled ligand or by a competitive binding assay, all of which are described more extensively below. HPR1 and HPR2 polypeptides having intracellular signaling activity bind ligand molecules when in association with other receptor polypeptides to form a homo- or heteromeric complex, with ligand binding initiating a signaling cascade. The intracellular signaling activity is associated with the cytoplasmic domain of certain HPR1 and HPR2 polypeptides. Thus, for uses requiring intracellular signaling activity, preferred HPR1 and HPR2 polypeptides include those having the cytoplasmic domain, and in particular having certain conserved domains (such as the Box 1 motif, the Trp residue at position 581 of SEQ ID NO:4, the Box 2 motif, the Asp-containing motifs between amino acids 588 and 639 of SEQ ID NO:4, or the Box 3 motif) and conserved cytoplasmic tyrosine residues, and exhibiting intracellular signaling biological activity. Preferred HPR1 and HPR2 polypeptides further include oligomers or fusion polypeptides comprising at least one cytoplasmic portion of one or more HPR1 and/or HPR2 polypeptides, and fragments of any of these polypeptides that have intracellular signaling activity. The intracellular signaling activity of HPR1 and HPR2 polypeptides may be determined, for example, through assays to detect phosphorylation of the HPR1 polypeptide, the HPR2 polypeptide, or downstream polypeptides in signaling cascades such as the JAK/STAT or ERK/MAPK pathways, or in assays that measure biological activities related to the signal transmission, such as stimulation or suppression of cell proliferation, differentiation, or activation. One example of an assay to measure cytokine-binding and cell-proliferation activity involves expressing a polypeptide of the invention in Ba/F3 cells, exposing the polypeptide-expressing cells to radioactively labeled cytokine, and measuring specific cytokine binding to cells and uptake of 3H-thymidine by cells in response to cytokine, as described in Presky et al., 1996, Proc Natl Acad Sci USA 93: 14002-14007. Further examples of such assays are described herein and in Ernst et al., 1999, J Biol Chem 274(14): 9729-9737. Soluble forms of hematopoietin receptors comprising one or more extracellular domains of the hematopoietin receptor, such as soluble forms of HPR1 and HPR2, may also be used in assays to measure their effect on cell growth, proliferation, differentiation, or activation; in such assays the cells are contacted with the soluble form of the receptor and their growth,

5 proliferation, differentiation, or activation is measured, for example by measuring the incorporation of radioactive thymidine or by microscopic examination of treated and untreated cells.

The terms "HPR1 polypeptide activity" and "HPR2 polypeptide activity," as used herein, include any one or more of the following: ligand-binding activity and intracellular signaling activity (which includes effects on cell growth, proliferation, differentiation, or activation), as well as the ex vivo and in vivo activities of HPR1 and HPR2 polypeptides. The degree to which HPR1 and HPR2 polypeptides and fragments and other derivatives of these polypeptides exhibit these activities can be determined by standard assay methods as disclosed herein; those of skill in the art will appreciate that other, similar types of assays can be used to measure HPR1 and HPR2 biological activities.

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Another aspect of the biological activity of HPR1 and HPR2 polypeptides is the ability of members of these polypeptide families to bind particular binding partners such as cytokines, other hematopoietin receptor polypeptides, and intracellular signaling polypeptides, with the cytokine receptor domain binding to cytokines and the intracellular signaling domain binding to intracellular signaling polypeptides such as members of the JAK and SHP polypeptide families. The term "binding partner," as used herein, includes ligands, receptors, substrates, antibodies, other hematopoietin receptor polypeptides, the same HPR1 or HPR2 polypeptide (in the case of homotypic interactions), and any other molecule that interacts with an HPR1 or an HPR2 polypeptide through contact or proximity between particular portions of the binding partner and the HPR1 or HPR2 polypeptide. Because the cytokine receptor domains of HPR1 and HPR2 polypeptides bind to cytokines, an HPR1 or HPR2 cytokine receptor domain when expressed as a separate fragment from the rest of an HPR1 or HPR2 polypeptide, or as a soluble polypeptide, fused for example to an immunoglobulin Fc domain, is expected to disrupt the binding of endogenous HPR1 and/or HPR2 polypeptides to their binding partners. By binding to one or more binding partners, the separate cytokine receptor domain polypeptide likely prevents binding by the native HPR1 and/or HPR2 polypeptide(s), and so acts in a dominant negative fashion to inhibit the biological activities mediated via binding of HPR1 and/or HPR2 polypeptides to cytokines. Assays for evaluating the biological activities and partner-binding properties of HPR1 and HPR2 polypeptides are described further herein.

HPR1 and HPR2 polypeptides are involved in cell proliferation, differentiation, or activation diseases or conditions, that share as a common feature ligand-binding activity in their etiology. More specifically, the following cell proliferation conditions are those that are known or are likely to involve the biological activities of HPR1 and/or HPR2 polypeptides: pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, osteoporosis resulting from a lack of bone-forming cells, leukemia, tumour metastasis, and osteoporosis resulting from an excess of bone-resorbing cells. In addition, the following metabolic conditions involving hematopoietin receptor ligands such as leptin are those that are known or are likely to involve the biological activities of HPR1 and/or HPR2 polypeptides: obesity, cachexia, wasting, and AIDS-related weight loss. Also, the following prolactin-related conditions are those that are known or are likely to involve the biological activities of HPR1 and/or HPR2 polypeptides: deficient mammary development, infertility, breast cancer, and

prolactinoma. Blocking or inhibiting the interactions between members of the HPR1 and HPR2 polypeptide families and their substrates, ligands, receptors, binding partners, and or other interacting polypeptides is an aspect of the invention and provides methods for treating or ameliorating these diseases and conditions through the use of inhibitors of HPR1 and/or HPR2 polypeptide activity. Examples of such inhibitors or antagonists are described in more detail below. For certain conditions involving too little HPR1 or HPR2 polypeptide activity, methods of treating or ameliorating these conditions comprise increasing the amount or activity of HPR1 or HPR2 polypeptides by providing isolated HPR1 or HPR2 polypeptides or active fragments or fusion polypeptides thereof, or by providing compounds (agonists) that activate endogenous or exogenous HPR1 or HPR2 polypeptides.

#### **HPR1 and HPR2 Polypeptides**

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An HPR1 polypeptide is a polypeptide that shares a sufficient degree of amino acid identity or similarity to the human HPR1 polypeptide of SEQ ID NO:4 or the murine HPR1 polypeptide of SEQ ID NO:12 to (A) be identified by those of skill in the art as a polypeptide likely to share particular structural domains and/or (B) have biological activities in common with the HPR1 polypeptides of SEQ ID NO:4 and SEQ ID NO:12 and/or (C) bind to antibodies that also specifically bind to other HPR1 polypeptides. An HPR2 polypeptide is a polypeptide that shares a sufficient degree of amino acid identity or similarity to the HPR2 polypeptides of SEQ ID NOs 21, 23, 25, and 27 to (A) be identified by those of skill in the art as a polypeptide likely to share particular structural domains and/or (B) have biological activities in common with the HPR2 polypeptides of SEQ ID NOs 21, 23, 25, and 27 and/or (C) bind to antibodies that also specifically bind to other HPR2 polypeptides. HPR1 and HPR2 polypeptides can be isolated from naturally occurring sources, or have the same structure as naturally occurring HPR1 or HPR2 polypeptides, or can be produced to have structures that differ from naturally occurring HPR1 or HPR2 polypeptides. Polypeptides derived from any HPR1 or HPR2 polypeptide by any type of alteration (for example, but not limited to, insertions, deletions, or substitutions of amino acids; changes in the state of glycosylation of the polypeptide; refolding or isomerization to change its three-dimensional structure or self-association state; and changes to its association with other polypeptides or molecules) are also HPR1 or HPR2 polypeptides, respectively. Therefore, the polypeptides provided by the invention include polypeptides characterized by amino acid sequences similar to those of the HPR1 and HPR2 polypeptides described herein, but into which modifications are naturally provided or deliberately engineered. A polypeptide that shares biological activities in common with members of the HPR1 and/or HPR2 polypeptide family is a polypeptide having HPR1 and/or HPR2 polypeptide activity. Examples of biological activities exhibited by HPR1 and/or HPR2 polypeptides include, without limitation, ligand-binding activity and intracellular signaling.

The present invention provides both full-length and mature forms of HPR1 and HPR2 polypeptides. Full-length polypeptides are those having the complete primary amino acid sequence of the polypeptide as initially translated. The amino acid sequences of full-length polypeptides can be obtained, for example, by translation of the complete open reading frame ("ORF") of a cDNA

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molecule. Several full-length polypeptides can be encoded by a single genetic locus if multiple mRNA forms are produced from that locus by alternative splicing or by the use of multiple translation initiation sites. The "mature form" of a polypeptide refers to a polypeptide that has undergone posttranslational processing steps such as cleavage of the signal sequence or proteolytic cleavage to remove a prodomain. Multiple mature forms of a particular full-length polypeptide may be produced, for example by cleavage of the signal sequence at multiple sites, or by differential regulation of proteases that cleave the polypeptide. The mature form(s) of such polypeptide can be obtained by expression, in a suitable mammalian cell or other host cell, of a nucleic acid molecule that encodes the full-length polypeptide. The sequence of the mature form of the polypeptide may also be determinable from the amino acid sequence of the full-length form, through identification of signal sequences or protease cleavage sites. The HPR1 and HPR2 polypeptides of the invention also include those that result from post-transcriptional or post-translational processing events such as alternate mRNA processing which can yield alternative splice forms of HPR1 or HPR2 such as a truncated but biologically active polypeptide or, for example, a naturally occurring soluble form of the polypeptide. Also encompassed within the invention are variations attributable to proteolysis such as differences in the N- or C-termini upon expression in different types of host cells, due to proteolytic removal of one or more terminal amino acids from the polypeptide (generally from 1-5 terminal amino acids).

The invention further includes HPR1 and HPR2 polypeptides with or without associated native-pattern glycosylation. Polypeptides expressed in yeast or mammalian expression systems (e.g., COS-1 or CHO cells) can be similar to or significantly different from a native polypeptide in molecular weight and glycosylation pattern, depending upon the choice of expression system. Expression of polypeptides of the invention in bacterial expression systems, such as E. coli, provides non-glycosylated molecules. Further, a given preparation can include multiple differentially glycosylated species of the polypeptide. Glycosyl groups can be removed through conventional methods, in particular those utilizing glycopeptidase. In general, glycosylated polypeptides of the invention can be incubated with a molar excess of glycopeptidase (Boehringer Mannheim).

Species homologues of HPR1 and HPR2 polypeptides and of nucleic acids encoding them are also provided by the present invention. As used herein, a "species homologue" is a polypeptide or nucleic acid with a different species of origin from that of a given polypeptide or nucleic acid, but with significant sequence similarity to the given polypeptide or nucleic acid, as determined by those of skill in the art. Species homologues can be isolated and identified by making suitable probes or primers from polynucleotides encoding the amino acid sequences provided herein and screening a suitable nucleic acid source from the desired species. The invention also encompasses allelic variants of HPR1 and HPR2 polypeptides and nucleic acids encoding them; that is, naturally-occurring alternative forms of such polypeptides and nucleic acids in which differences in amino acid or nucleotide sequence are attributable to genetic polymorphism (allelic variation among individuals within a population).

Fragments of the HPR1 and HPR2 polypeptides of the present invention are encompassed by the present invention and can be in linear form or cyclized using known methods, for example, as

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described in Saragovi, et al., Bio/Technology 10, 773-778 (1992) and in McDowell, et al., J. Amer. Chem. Soc. 114 9245-9253 (1992). Polypeptides and polypeptide fragments of the present invention, and nucleic acids encoding them, include polypeptides and nucleic acids with amino acid or nucleotide sequence lengths that are at least 25% (more preferably at least 50%, or at least 60%, or at least 70%, and most preferably at least 80%) of the length of an HPR1 polypeptide or of an HPR2 polypeptide, and have at least 60% sequence identity (more preferably at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, or at least 99%, and most preferably at least 99.5%) with that HPR1 or HPR2 polypeptide or encoding nucleic acid, where sequence identity is determined by comparing the amino acid sequences of the polypeptides when aligned so as to maximize overlap and identity while minimizing sequence gaps. Also included in the present invention are polypeptides and polypeptide fragments, and nucleic acids encoding them, that contain or encode a segment preferably comprising at least 8, or at least 10, or preferably at least 15, or more preferably at least 20, or still more preferably at least 30, or most preferably at least 40 contiguous amino acids. Such polypeptides and polypeptide fragments may also contain a segment that shares at least 70% sequence identity (more preferably at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, or at least 99%, and most preferably at least 99.5%) with any such segment of any of the HPR1 or HPR2 polypeptides, where sequence identity is determined by comparing the amino acid sequences of the polypeptides when aligned so as to maximize overlap and identity while minimizing sequence gaps. The percent identity can be determined by visual inspection and mathematical calculation. Preferably, the comparison is done using a computer program. An exemplary, preferred computer program is the Genetics Computer Group (GCG; Madison, WI) Wisconsin package version 10.0 program, 'GAP.' The preferred default parameters for the 'GAP' program includes: (1) The GCG implementation of comparison matrices for nucleotides and amino acids; such as a unary comparison matrix (containing a value of 1 for identities and 0 for non-identities) for nucleotides, and the weighted comparison matrix of Gribskov and Burgess, Nucl. Acids Res. 14:6745, 1986, as described by Schwartz and Dayhoff, eds., Atlas of Polypeptide Sequence and Structure, National Biomedical Research Foundation, pp. 353-358, 1979; (2) a penalty of 30 for each gap and an additional penalty of 1 for each symbol in each gap for amino acid sequences, or penalty of 50 for each gap and an additional penalty of 3 for each symbol in each gap for nucleotide sequences; (3) no penalty for end gaps; and (4) no maximum penalty for long gaps. Another program useful for determining percent identify is the BESTFIT program, also available from the University of Wisconsin as part of the GCG computer package. Default parameters for using the BESTFIT program are the same as those described above for using the GAP program. Other programs used by those skilled in the art of sequence comparison can also be used, such as, for example, the UW-BLAST 2.0 algorithm or the BLASTN program version 2.0.9, available for use via the National Library of Medicine website: ncbi.nlm.nih.gov/gorf/wblast2.cgi. Standard default parameter settings for UW-BLAST 2.0 are described at the following Internet site: blast.wustl.edu/blast/README.html#References. In addition, the BLAST algorithm uses the BLOSUM62 amino acid scoring matix, and optional parameters that can

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be used are as follows: (A) inclusion of a filter to mask segments of the query sequence that have low compositional complexity (as determined by the SEG program of Wootton and Federhen (Computers and Chemistry, 1993); also see Wootton and Federhen, 1996, Analysis of compositionally biased regions in sequence databases, *Methods Enzymol.* 266: 554-71) or segments consisting of short-periodicity internal repeats (as determined by the XNU program of Claverie and States (Computers and Chemistry, 1993)), and (B) a statistical significance threshold for reporting matches against database sequences, or E-score (the expected probability of matches being found merely by chance, according to the stochastic model of Karlin and Altschul (1990); if the statistical significance ascribed to a match is greater than this E-score threshold, the match will not be reported.); preferred E-score threshold values are 0.5, or in order of increasing preference, 0.25, 0.1, 0.01, 0.001, 0.0001, 1e-10, 1e-20, 1e-30, 1e-40, 1e-50, 1e-75, or 1e-100.

"An isolated polypeptide consisting essentially of an amino acid sequence" means that the polypeptide may have, in addition to said amino acid sequence, additional material covalently linked to either or both ends of the polypeptide, said additional material preferably between 1 and 10,000 additional amino acids covalently linked to either end, each end, or both ends of polypeptide, and more preferably between 1 and 1,000 additional amino acids covalently linked to either end, each end, or both ends of the polypeptide, and most preferably between 1 and 100 additional amino acids covalently linked to either end, each end, or both ends of the polypeptide. In preferred embodiments, covalent linkage of additional amino acids to either end, each end, or both ends of the polypeptide results in a novel combined amino acid sequence that is neither naturally occurring nor disclosed in the art.

The present invention also provides for soluble forms of HPR1 and HPR2 polypeptides comprising or consisting essentially of certain fragments or domains of these polypeptides, and particularly those comprising the extracellular domain or one or more fragments of the extracellular domain. Soluble polypeptides are polypeptides that are capable of being secreted from the cells in which they are expressed. In such forms part or all of the intracellular and transmembrane domains of the polypeptide are deleted such that the polypeptide is fully secreted from the cell in which it is expressed. The intracellular and transmembrane domains of polypeptides of the invention can be identified in accordance with known techniques for determination of such domains from sequence information. Soluble HPR1 and HPR2 polypeptides also include those polypeptides which include part of the transmembrane region, provided that the soluble HPR1 or HPR2 polypeptide is capable of being secreted from a cell, and preferably retains HPR1 and/or HPR2 polypeptide activity. Soluble HPR1 and HPR2 polypeptides further include oligomers or fusion polypeptides comprising the extracellular portion of at least one HPR1 or HPR2 polypeptide, and fragments of any of these polypeptides that have HPR1 and/or HPR2 polypeptide activity. A secreted soluble polypeptide can be identified (and distinguished from its non-soluble membrane-bound counterparts) by separating intact cells which express the desired polypeptide from the culture medium, e.g., by centrifugation, and assaying the medium (supernatant) for the presence of the desired polypeptide. The presence of the desired polypeptide in the medium indicates that the polypeptide was secreted from the cells and thus is a

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soluble form of the polypeptide. The use of soluble forms of HPR1 or HPR2 polypeptides is advantageous for many applications. Purification of the polypeptides from recombinant host cells is facilitated, since the soluble polypeptides are secreted from the cells. Moreover, soluble polypeptides are generally more suitable than membrane-bound forms for parenteral administration and for many enzymatic procedures.

In another aspect of the invention, preferred polypeptides comprise various combinations of HPR1 and/or HPR2 polypeptide domains, such as the cytokine receptor domain and the intracellular signaling domain. Accordingly, polypeptides of the present invention and nucleic acids encoding them include those comprising or encoding two or more copies of a domain such as the cytokine receptor domain, two or more copies of a domain such as the intracellular signaling domain, or at least one copy of each domain, and these domains can be presented in any order within such polypeptides.

Further modifications in the peptide or DNA sequences can be made by those skilled in the art using known techniques. Modifications of interest in the polypeptide sequences can include the alteration, substitution, replacement, insertion or deletion of a selected amino acid. For example, one or more of the cysteine residues can be deleted or replaced with another amino acid to alter the conformation of the molecule, an alteration which may involve preventing formation of incorrect intramolecular disulfide bridges upon folding or renaturation. Techniques for such alteration, substitution, replacement, insertion or deletion are well known to those skilled in the art (see, e.g., U.S. Pat. No. 4,518,584). As another example, N-glycosylation sites in the polypeptide extracellular domain can be modified to preclude glycosylation, allowing expression of a reduced carbohydrate analog in mammalian and yeast expression systems. N-glycosylation sites in eukaryotic polypeptides are characterized by an amino acid triplet Asn-X-Y, wherein X is any amino acid except Pro and Y is Ser or Thr. Appropriate substitutions, additions, or deletions to the nucleotide sequence encoding these triplets will result in prevention of attachment of carbohydrate residues at the Asn side chain. Alteration of a single nucleotide, chosen so that Asn is replaced by a different amino acid, for example, is sufficient to inactivate an N-glycosylation site. Alternatively, the Ser or Thr can by replaced with another amino acid, such as Ala. Known procedures for inactivating N-glycosylation sites in polypeptides include those described in U.S. Patent 5,071,972 and EP 276,846. Additional variants within the scope of the invention include polypeptides that can be modified to create derivatives thereof by forming covalent or aggregative conjugates with other chemical moieties, such as glycosyl groups, lipids, phosphate, acetyl groups and the like. Covalent derivatives can be prepared by linking the chemical moieties to functional groups on amino acid side chains or at the N-terminus or C-terminus of a polypeptide. Conjugates comprising diagnostic (detectable) or therapeutic agents attached thereto are contemplated herein. Preferably, such alteration, substitution, replacement, insertion or deletion retains the desired activity of the polypeptide or a substantial equivalent thereof. One example is a variant that binds with essentially the same binding affinity as does the native form. Binding affinity can be measured by conventional procedures, e.g., as described in U.S. Patent No. 5,512,457 and as set forth herein.

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Other derivatives include covalent or aggregative conjugates of the polypeptides with other polypeptides or polypeptides, such as by synthesis in recombinant culture as N-terminal or C-terminal fusions. Examples of fusion polypeptides are discussed below in connection with oligomers. Further, fusion polypeptides can comprise peptides added to facilitate purification and identification. Such peptides include, for example, poly-His or the antigenic identification peptides described in U.S. Patent No. 5,011,912 and in Hopp et al., *Bio/Technology* 6:1204, 1988. One such peptide is the FLAG® peptide, which is highly antigenic and provides an epitope reversibly bound by a specific monoclonal antibody, enabling rapid assay and facile purification of expressed recombinant polypeptide. A murine hybridoma designated 4E11 produces a monoclonal antibody that binds the FLAG® peptide in the presence of certain divalent metal cations, as described in U.S. Patent 5,011,912. The 4E11 hybridoma cell line has been deposited with the American Type Culture Collection under accession no. HB 9259. Monoclonal antibodies that bind the FLAG® peptide are available from Eastman Kodak Co., Scientific Imaging Systems Division, New Haven, Connecticut.

Encompassed by the invention are oligomers or fusion polypeptides that contain an HPR1 polypeptide and/or an HPR2 polypeptide, one or more fragments of HPR1 and/or HPR2 polypeptides, or any of the derivative or variant forms of HPR1 and HPR2 polypeptides as disclosed herein. In particular embodiments, the oligomers comprise soluble HPR1 and/or HPR2 polypeptides. Oligomers can be in the form of covalently linked or non-covalently-linked multimers, including dimers, trimers, or higher oligomers. In one aspect of the invention, the oligomers maintain the binding ability of the polypeptide components and provide therefor, bivalent, trivalent, etc., binding sites. In an alternative embodiment the invention is directed to oligomers comprising multiple HPR1 and/or HPR2 polypeptides joined via covalent or non-covalent interactions between peptide moieties fused to the polypeptides, such peptides having the property of promoting oligomerization. Leucine zippers and certain polypeptides derived from antibodies are among the peptides that can promote oligomerization of the polypeptides attached thereto, as described in more detail below.

In embodiments where variants of the HPR1 and/or HPR2 polypeptides are constructed to include a membrane-spanning domain, they will form a Type I membrane polypeptide. Membrane-spanning HPR1 and/or HPR2 polypeptides can be fused with extracellular domains of receptor polypeptides for which the ligand is known. Such fusion polypeptides can then be manipulated to control the intracellular signaling pathways triggered by the membrane-spanning HPR1 or HPR2 polypeptide. HPR1 and HPR2 polypeptides that span the cell membrane can also be fused with agonists or antagonists of cell-surface receptors, or cellular adhesion molecules to further modulate HPR1 and/or HPR2 intracellular effects. In another aspect of the present invention, interleukins can be situated between the preferred HPR1 or HPR2 polypeptide fragment and other fusion polypeptide domains.

<u>Immunoglobulin-based Oligomers</u>. The polypeptides of the invention or fragments thereof can be fused to molecules such as immunoglobulins for many purposes, including increasing the valency of polypeptide binding sites. For example, fragments of an HPR1 polypeptide and/or of an

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HPR2 polypeptide can be fused directly or through linker sequences to the Fc portion of an immunoglobulin. For a bivalent form of the polypeptide, such a fusion could be to the Fc portion of an IgG molecule. Other immunoglobulin isotypes can also be used to generate such fusions. For example, a polypeptide-IgM fusion would generate a decayalent form of the polypeptide of the invention. The term "Fc polypeptide" as used herein includes native and mutein forms of polypeptides 10 made up of the Fc region of an antibody comprising any or all of the CH domains of the Fc region. Truncated forms of such polypeptides containing the hinge region that promotes dimerization are also included. Preferred Fc polypeptides comprise an Fc polypeptide derived from a human IgG1 antibody. As one alternative, an oligomer is prepared using polypeptides derived from immunoglobulins. Preparation of fusion polypeptides comprising certain heterologous polypeptides fused to various 15 portions of antibody-derived polypeptides (including the Fc domain) has been described, e.g., by Ashkenazi et al. (PNAS USA 88:10535, 1991); Byrn et al. (Nature 344:677, 1990); and Hollenbaugh and Aruffo ("Construction of Immunoglobulin Fusion Polypeptides", in Current Protocols in Immunology, Suppl. 4, pages 10.19.1 - 10.19.11, 1992). Methods for preparation and use of immunoglobulin-based oligomers are well known in the art. One embodiment of the present invention 20 is directed to a dimer comprising two fusion polypeptides created by fusing a polypeptide of the invention to an Fc polypeptide derived from an antibody. A gene fusion encoding the polypeptide/Fc fusion polypeptide is inserted into an appropriate expression vector. Polypeptide/Fc fusion polypeptides are expressed in host cells transformed with the recombinant expression vector, and allowed to assemble much like antibody molecules, whereupon interchain disulfide bonds form 25 between the Fc moieties to yield divalent molecules. One suitable Fc polypeptide, described in PCT application WO 93/10151, is a single chain polypeptide extending from the N-terminal hinge region to the native C-terminus of the Fc region of a human IgG1 antibody. Another useful Fc polypeptide is the Fc mutein described in U.S. Patent 5,457,035 and in Baum et al., (EMBO J. 13:3992-4001, 1994). The amino acid sequence of this mutein is identical to that of the native Fc sequence presented in WO 30 93/10151, except that amino acid 19 has been changed from Leu to Ala, amino acid 20 has been changed from Leu to Glu, and amino acid 22 has been changed from Gly to Ala. The mutein exhibits reduced affinity for Fc receptors. The above-described fusion polypeptides comprising Fc moieties (and oligomers formed therefrom) offer the advantage of facile purification by affinity chromatography over Polypeptide A or Polypeptide G columns. In other embodiments, the polypeptides of the 35 invention can be substituted for the variable portion of an antibody heavy or light chain. If fusion polypeptides are made with both heavy and light chains of an antibody, it is possible to form an oligomer with as many as four HPR1 and/or HPR2 extracellular regions.

Peptide-linker Based Oligomers. Alternatively, the oligomer is a fusion polypeptide comprising multiple HPR1 and/or HPR2 polypeptides, with or without peptide linkers (spacer peptides). Among the suitable peptide linkers are those described in U.S. Patents 4,751,180 and 4,935,233. A DNA sequence encoding a desired peptide linker can be inserted between, and in the same reading frame as, the DNA sequences of the invention, using any suitable conventional technique.

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For example, a chemically synthesized oligonucleotide encoding the linker can be ligated between the sequences. In particular embodiments, a fusion polypeptide comprises from two to four soluble HPR1 and/or HPR2 polypeptides, separated by peptide linkers. Suitable peptide linkers, their combination with other polypeptides, and their use are well known by those skilled in the art.

Leucine-Zippers. Another method for preparing the oligomers of the invention involves use of a leucine zipper. Leucine zipper domains are peptides that promote oligomerization of the polypeptides in which they are found. Leucine zippers were originally identified in several DNA-binding polypeptides (Landschulz et al., *Science* 240:1759, 1988), and have since been found in a variety of different polypeptides. Among the known leucine zippers are naturally occurring peptides and derivatives thereof that dimerize or trimerize. The zipper domain (also referred to herein as an oligomerizing, or oligomer-forming, domain) comprises a repetitive heptad repeat, often with four or five leucine residues interspersed with other amino acids. Use of leucine zippers and preparation of oligomers using leucine zippers are well known in the art.

Other fragments and derivatives of the sequences of polypeptides which would be expected to retain polypeptide activity in whole or in part and may thus be useful for screening or other immunological methodologies can also be made by those skilled in the art given the disclosures herein. Such modifications are believed to be encompassed by the present invention.

#### Nucleic Acids Encoding HPR1 Polypeptides and Nucleic Acids Encoding HPR2 Polypeptides

Encompassed within the invention are nucleic acids encoding HPR1 polypeptides and nucleic acids encoding HPR2 polypeptides. These nucleic acids can be identified in several ways, including isolation of genomic or cDNA molecules from a suitable source. Nucleotide sequences corresponding to the amino acid sequences described herein, to be used as probes or primers for the isolation of nucleic acids or as query sequences for database searches, can be obtained by "back-translation" from the amino acid sequences, or by identification of regions of amino acid identity with polypeptides for which the coding DNA sequence has been identified. The well-known polymerase chain reaction (PCR) procedure can be employed to isolate and amplify a DNA sequence encoding an HPR1 or HPR2 polypeptide or a desired combination of HPR1 and/or HPR2 polypeptide fragments. Oligonucleotides that define the desired termini of the combination of DNA fragments are employed as 5' and 3' primers. The oligonucleotides can additionally contain recognition sites for restriction endonucleases, to facilitate insertion of the amplified combination of DNA fragments into an expression vector. PCR techniques are described in Saiki et al., Science 239:487 (1988); Recombinant DNA Methodology, Wu et al., eds., Academic Press, Inc., San Diego (1989), pp. 189-196; and PCR Protocols: A Guide to Methods and Applications, Innis et. al., eds., Academic Press, Inc. (1990).

Nucleic acid molecules of the invention include DNA and RNA in both single-stranded and double-stranded form, as well as the corresponding complementary sequences. DNA includes, for example, cDNA, genomic DNA, chemically synthesized DNA, DNA amplified by PCR, and combinations thereof. The nucleic acid molecules of the invention include full-length genes or cDNA

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molecules as well as a combination of fragments thereof. The nucleic acids of the invention are preferentially derived from human sources, but the invention includes those derived from non-human species, as well.

An "isolated nucleic acid" is a nucleic acid that has been separated from adjacent genetic sequences present in the genome of the organism from which the nucleic acid was isolated, in the case of nucleic acids isolated from naturally-occurring sources. In the case of nucleic acids synthesized enzymatically from a template or chemically, such as PCR products, cDNA molecules, or oligonucleotides for example, it is understood that the nucleic acids resulting from such processes are isolated nucleic acids. An isolated nucleic acid molecule refers to a nucleic acid molecule in the form of a separate fragment or as a component of a larger nucleic acid construct. In one preferred embodiment, the invention relates to certain isolated nucleic acids that are substantially free from contaminating endogenous material. The nucleic acid molecule has preferably been derived from DNA or RNA isolated at least once in substantially pure form and in a quantity or concentration enabling identification, manipulation, and recovery of its component nucleotide sequences by standard biochemical methods (such as those outlined in Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, NY (1989)). Such sequences are preferably provided and/or constructed in the form of an open reading frame uninterrupted by internal non-translated sequences, or introns, that are typically present in eukaryotic genes. Sequences of non-translated DNA can be present 5' or 3' from an open reading frame, where the same do not interfere with manipulation or expression of the coding region.

"An isolated nucleic acid consisting essentially of a nucleotide sequence" means that the nucleic acid may have, in addition to said nucleotide sequence, additional material covalently linked to either or both ends of the nucleic acid molecule, said additional material preferably between 1 and 100,000 additional nucleotides covalently linked to either end, each end, or both ends of the nucleic acid molecule, and more preferably between 1 and 1,000 additional nucleotides covalently linked to either end, each end, or both ends of the nucleic acid molecule, and most preferably between 10 and 100 additional nucleotides covalently linked to either end, each end, or both ends of the nucleic acid molecule. In preferred embodiments, covalent linkage of additional nucleotides to either end, each end, or both ends of the nucleic acid molecule results in a novel combined nucleotide sequence that is neither naturally occurring nor disclosed in the art. An isolated nucleic acid consisting essentially of a nucleotide sequence may be an expression vector or other construct comprising said nucleotide sequence.

The present invention also includes nucleic acids that hybridize under moderately stringent conditions, and more preferably highly stringent conditions, to nucleic acids encoding HPR1 polypeptides and/or nucleic acids encoding HPR2 polypeptides described herein. The basic parameters affecting the choice of hybridization conditions and guidance for devising suitable conditions are set forth by Sambrook,, Fritsch, and Maniatis (1989, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., chapters 9 and 11; and Current Protocols

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in Molecular Biology, 1995, Ausubel et al., eds., John Wiley & Sons, Inc., sections 2.10 and 6.3-6.4), and can be readily determined by those having ordinary skill in the art based on, for example, the length and/or base composition of the DNA. One way of achieving moderately stringent conditions involves the use of a prewashing solution containing 5 x SSC, 0.5% SDS, 1.0 mM EDTA (pH 8.0), hybridization buffer of about 50% formamide, 6 x SSC, and a hybridization temperature of about 55 degrees C (or other similar hybridization solutions, such as one containing about 50% formamide, with a hybridization temperature of about 42 degrees C), and washing conditions of about 60 degrees C, in 0.5 x SSC, 0.1% SDS. Generally, highly stringent conditions are defined as hybridization conditions as above, but with washing at approximately 68 degrees C, 0.2 x SSC, 0.1% SDS. SSPE (1xSSPE is 0.15M NaCl, 10 mM NaH.sub.2 PO.sub.4, and 1.25 mM EDTA, pH 7.4) can be substituted for SSC (1xSSC is 0.15M NaCl and 15 mM sodium citrate) in the hybridization and wash buffers; washes are performed for 15 minutes after hybridization is complete. It should be understood that the wash temperature and wash salt concentration can be adjusted as necessary to achieve a desired degree of stringency by applying the basic principles that govern hybridization reactions and duplex stability, as known to those skilled in the art and described further below (see, e.g., Sambrook et al., 1989). When hybridizing a nucleic acid to a target nucleic acid of unknown sequence, the hybrid length is assumed to be that of the hybridizing nucleic acid. When nucleic acids of known sequence are hybridized, the hybrid length can be determined by aligning the sequences of the nucleic acids and identifying the region or regions of optimal sequence complementarity. The hybridization temperature for hybrids anticipated to be less than 50 base pairs in length should be 5 to 10.degrees C less than the melting temperature (Tm) of the hybrid, where Tm is determined according to the following equations. For hybrids less than 18 base pairs in length, Tm (degrees C) = 2(# of A + T bases) + 4(# of #G + C bases). For hybrids above 18 base pairs in length, Tm (degrees C) =  $81.5 + 16.6(\log_{10} [\text{Na}^+]) + 0.41(\% \text{ G} + \text{C})$ - (600/N), where N is the number of bases in the hybrid, and [Na<sup>+</sup>] is the concentration of sodium ions in the hybridization buffer ([Na<sup>+</sup>] for 1xSSC = 0.165M). Preferably, each such hybridizing nucleic acid has a length that is at least 15 nucleotides (or more preferably at least 18 nucleotides, or at least 20 nucleotides, or at least 25 nucleotides, or at least 30 nucleotides, or at least 40 nucleotides, or most preferably at least 50 nucleotides), or at least 25% (more preferably at least 50%, or at least 60%, or at least 70%, and most preferably at least 80%) of the length of the nucleic acid of the present invention to which it hybridizes, and has at least 60% sequence identity (more preferably at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, or at least 99%, and most preferably at least 99.5%) with the nucleic acid of the present invention to which it hybridizes, where sequence identity is determined by comparing the sequences of the hybridizing nucleic acids when aligned so as to maximize overlap and identity while minimizing sequence gaps as described in more detail above.

The present invention also provides genes corresponding to the nucleic acid sequences disclosed herein. "Corresponding genes" or "corresponding genomic nucleic acids" are the regions of the genome that are transcribed to produce the mRNAs from which cDNA nucleic acid sequences are

derived and can include contiguous regions of the genome necessary for the regulated expression of such genes. Corresponding genes can therefore include but are not limited to coding sequences, 5' and 3' untranslated regions, alternatively spliced exons, introns, promoters, enhancers, and silencer or suppressor elements. Corresponding genomic nucleic acids can include 10000 basepairs (more preferably, 5000 basepairs, still more preferably, 2500 basepairs, and most preferably, 1000 basepairs) of genomic nucleic acid sequence upstream of the first nucleotide of the genomic sequence corresponding to the initiation codon of the HPR1 coding sequence or of the HPR2 coding sequence, and 10000 basepairs (more preferably, 5000 basepairs, still more preferably, 2500 basepairs, and most preferably, 1000 basepairs) of genomic nucleic acid sequence downstream of the last nucleotide of the genomic sequence corresponding to the termination codon of the HPR1 coding sequence or of the HPR2 coding sequence. The corresponding genes or genomic nucleic acids can be isolated in accordance with known methods using the sequence information disclosed herein. Such methods include the preparation of probes or primers from the disclosed sequence information for identification and/or amplification of genes in appropriate genomic libraries or other sources of genomic materials. An "isolated gene" or an "isolated genomic nucleic acid" is a genomic nucleic acid that has been separated from the adjacent genomic sequences present in the genome of the organism from which the genomic nucleic acid was isolated.

#### Methods for Making and Purifying HPR1 and HPR2 Polypeptides

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Methods for making HPR1 and HPR2 polypeptides are described below. Expression, isolation, and purification of the polypeptides and fragments of the invention can be accomplished by any suitable technique, including but not limited to the following methods. The isolated nucleic acid of the invention can be operably linked to an expression control sequence such as the pDC409 vector (Giri et al., 1990, EMBO J., 13: 2821) or the derivative pDC412 vector (Wiley et al., 1995, Immunity 3: 673). The pDC400 series vectors are useful for transient mammalian expression systems, such as CV-1 or 293 cells. Alternatively, the isolated nucleic acid of the invention can be linked to expression vectors such as pDC312, pDC316, or pDC317 vectors, which are useful for stable mammalian expression systems, such as CHO cells or their derivatives. Other expression control sequences and cloning technologies can also be used to produce the polypeptide recombinantly, such as the pMT2 or pED expression vectors (Kaufman et al., 1991, Nucleic Acids Res. 19: 4485-4490; and Pouwels et al., 1985, Cloning Vectors: A Laboratory Manual, Elsevier, New York) and the GATEWAY Vectors (lifetech.com/Content/Tech-Online/molecular\_biology/manuals\_pps/11797016.pdf; Life Technologies; Rockville, MD). Many suitable expression control sequences are known in the art. General methods of expressing recombinant polypeptides are also known and are exemplified in R. Kaufman, Methods in Enzymology 185, 537-566 (1990). As used herein "operably linked" means that the nucleic acid of the invention and an expression control sequence are situated within a construct, vector, or cell in such a way that the polypeptide encoded by the nucleic acid is expressed when appropriate molecules (such as polymerases) are present. As one embodiment of the invention, at least one expression control

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sequence is operably linked to the nucleic acid of the invention in a recombinant host cell or progeny thereof, the nucleic acid and/or expression control sequence having been introduced into the host cell by transformation or transfection, for example, or by any other suitable method. As another embodiment of the invention, at least one expression control sequence is integrated into the genome of a recombinant host cell such that it is operably linked to a nucleic acid sequence encoding a polypeptide of the invention. In a further embodiment of the invention, at least one expression control sequence is operably linked to a nucleic acid of the invention through the action of a trans-acting factor such as a transcription factor, either *in vitro* or in a recombinant host cell.

In addition, a sequence encoding an appropriate signal peptide (native or heterologous) can be incorporated into expression vectors. The choice of signal peptide or leader can depend on factors such as the type of host cells in which the recombinant polypeptide is to be produced. To illustrate, examples of heterologous signal peptides that are functional in mammalian host cells include the signal sequence for interleukin-7 (IL-7) described in United States Patent 4,965,195; the signal sequence for interleukin-2 receptor described in Cosman et al., Nature 312:768 (1984); the interleukin-4 receptor signal peptide described in EP 367,566; the type I interleukin-1 receptor signal peptide described in U.S. Patent 4,968,607; and the type II interleukin-1 receptor signal peptide described in EP 460,846. A DNA sequence for a signal peptide (secretory leader) can be fused in frame to the nucleic acid sequence of the invention so that the DNA is initially transcribed, and the mRNA translated, into a fusion polypeptide comprising the signal peptide. A signal peptide that is functional in the intended host cells promotes extracellular secretion of the polypeptide. The signal peptide is cleaved from the polypeptide upon secretion of polypeptide from the cell. The skilled artisan will also recognize that the position(s) at which the signal peptide is cleaved can differ from that predicted by computer program, and can vary according to such factors as the type of host cells employed in expressing a recombinant polypeptide. A polypeptide preparation can include a mixture of polypeptide molecules having different N-terminal amino acids, resulting from cleavage of the signal peptide at more than one site.

Established methods for introducing DNA into mammalian cells have been described (Kaufman, R.J., Large Scale Mammalian Cell Culture, 1990, pp. 15-69). Additional protocols using commercially available reagents, such as Lipofectamine lipid reagent (Gibco/BRL) or Lipofectamine-Plus lipid reagent, can be used to transfect cells (Felgner et al., Proc. Natl. Acad. Sci. USA 84:7413-7417, 1987). In addition, electroporation can be used to transfect mammalian cells using conventional procedures, such as those in Sambrook et al. (Molecular Cloning: A Laboratory Manual, 2 ed. Vol. 1-3, Cold Spring Harbor Laboratory Press, 1989). Selection of stable transformants can be performed using methods known in the art, such as, for example, resistance to cytotoxic drugs. Kaufman et al., Meth. in Enzymology 185:487-511, 1990, describes several selection schemes, such as dihydrofolate reductase (DHFR) resistance. A suitable strain for DHFR selection can be CHO strain DX-B11, which is deficient in DHFR (Urlaub and Chasin, Proc. Natl. Acad. Sci. USA 77:4216-4220, 1980). Other examples of selectable markers that can be incorporated into an expression vector include cDNAs

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conferring resistance to antibiotics, such as G418 and hygromycin B. Cells harboring the vector can be selected on the basis of resistance to these compounds.

Alternatively, gene products can be obtained via homologous recombination, or "gene targeting," techniques. Such techniques employ the introduction of exogenous transcription control elements (such as the CMV promoter or the like) in a particular predetermined site on the genome, to induce expression of the endogenous nucleic acid sequence of interest (see, for example, U.S. Patent No. 5,272,071). The location of integration into a host chromosome or genome can be easily determined by one of skill in the art, given the known location and sequence of the gene. In a preferred embodiment, the present invention also contemplates the introduction of exogenous transcriptional control elements in conjunction with an amplifiable gene, to produce increased amounts of the gene product, again, without the need for isolation of the gene sequence itself from the host cell.

A number of types of cells can act as suitable host cells for expression of the polypeptide. Mammalian host cells include, for example, the COS-7 line of monkey kidney cells (ATCC CRL 1651) (Gluzman et al., Cell 23:175, 1981), L cells, C127 cells, 3T3 cells (ATCC CCL 163), Chinese hamster ovary (CHO) cells, HeLa cells, BHK (ATCC CRL 10) cell lines, the CV1/EBNA cell line derived from the African green monkey kidney cell line CV1 (ATCC CCL 70) as described by McMahan et al. (EMBO J. 10: 2821, 1991), human kidney 293 cells, human epidermal A431 cells, human Colo205 cells, other transformed primate cell lines, normal diploid cells, cell strains derived from in vitro culture of primary tissue, primary explants, HL-60, U937, HaK or Jurkat cells. Alternatively, it is possible to produce the polypeptide in lower eukaryotes such as yeast or in prokaryotes such as bacteria. Potentially suitable yeasts include Saccharomyces cerevisiae, Schizosaccharomyces pombe, Kluyveromyces strains, Candida, or any yeast strain capable of expressing heterologous polypeptides. Potentially suitable bacterial strains include Escherichia coli, Bacillus subtilis, Salmonella typhimurium, or any bacterial strain capable of expressing heterologous polypeptides. If the polypeptide is made in yeast or bacteria, it may be necessary to modify the polypeptide produced therein, for example by phosphorylation or glycosylation of the appropriate sites, in order to obtain the functional polypeptide. Such covalent attachments can be accomplished using known chemical or enzymatic methods. The polypeptide can also be produced by operably linking the isolated nucleic acid of the invention to suitable control sequences in one or more insect expression vectors, and employing an insect expression system. Materials and methods for baculovirus/insect cell expression systems are commercially available in kit form from, e.g., Invitrogen, San Diego, Calif., U.S.A. (the MaxBac® kit), and such methods are well known in the art, as described in Summers and Smith, Texas Agricultural Experiment Station Bulletin No. 1555 (1987), and Luckow and Summers, Bio/Technology 6:47 (1988). Cell-free translation systems could also be employed to produce polypeptides using RNAs derived from nucleic acid constructs disclosed herein. A host cell that comprises an isolated nucleic acid of the invention, preferably operably linked to at least one expression control sequence, is a "recombinant host cell".

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The polypeptide of the invention can be prepared by culturing transformed host cells under culture conditions suitable to express the recombinant polypeptide. The resulting expressed polypeptide can then be purified from such culture (i.e., from culture medium or cell extracts) using known purification processes, such as gel filtration and ion exchange chromatography. The purification of the polypeptide can also include an affinity column containing agents which will bind to the polypeptide; one or more column steps over such affinity resins as concanavalin A-agarose, heparintoyopearl® or Cibacrom blue 3GA Sepharose®; one or more steps involving hydrophobic interaction chromatography using such resins as phenyl ether, butyl ether, or propyl ether; or immunoaffinity chromatography. Alternatively, the polypeptide of the invention can also be expressed in a form which will facilitate purification. For example, it can be expressed as a fusion polypeptide, such as those of maltose binding polypeptide (MBP), glutathione-S-transferase (GST) or thioredoxin (TRX). Kits for expression and purification of such fusion polypeptides are commercially available from New England BioLab (Beverly, Mass.), Pharmacia (Piscataway, N.J.) and InVitrogen, respectively. The polypeptide can also be tagged with an epitope and subsequently purified by using a specific antibody directed to such epitope. One such epitope (FLAG®) is commercially available from Kodak (New Haven, Conn.). Finally, one or more reverse-phase high performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media, e.g., silica gel having pendant methyl or other aliphatic groups, can be employed to further purify the polypeptide. Some or all of the foregoing purification steps, in various combinations, can also be employed to provide a substantially homogeneous isolated recombinant polypeptide. The polypeptide thus purified is substantially free of other mammalian polypeptides and is defined in accordance with the present invention as an "isolated polypeptide"; such isolated polypeptides of the invention include isolated antibodies that bind to HPR1 and/or HPR2 polypeptides, fragments, variants, binding partners etc. The polypeptide of the invention can also be expressed as a product of transgenic animals, e.g., as a component of the milk of transgenic cows, goats, pigs, or sheep which are characterized by somatic or germ cells containing a nucleotide sequence encoding the polypeptide.

It is also possible to utilize an affinity column comprising a polypeptide-binding polypeptide of the invention, such as a monoclonal antibody generated against polypeptides of the invention, to affinity-purify expressed polypeptides. These polypeptides can be removed from an affinity column using conventional techniques, e.g., in a high salt elution buffer and then dialyzed into a lower salt buffer for use or by changing pH or other components depending on the affinity matrix utilized, or be competitively removed using the naturally occurring substrate of the affinity moiety, such as a polypeptide derived from the invention. In this aspect of the invention, polypeptide-binding polypeptides, such as the anti-polypeptide antibodies of the invention or other polypeptides that can interact with the polypeptide of the invention, can be bound to a solid phase support such as a column chromatography matrix or a similar substrate suitable for identifying, separating, or purifying cells that express polypeptides of the invention on their surface. Adherence of polypeptide-binding polypeptides of the invention to a solid phase contacting surface can be accomplished by any means, for example,

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magnetic microspheres can be coated with these polypeptide-binding polypeptides and held in the incubation vessel through a magnetic field. Suspensions of cell mixtures are contacted with the solid phase that has such polypeptide-binding polypeptides thereon. Cells having polypeptides of the invention on their surface bind to the fixed polypeptide-binding polypeptide and unbound cells then are washed away. This affinity-binding method is useful for purifying, screening, or separating such polypeptide-expressing cells from solution. Methods of releasing positively selected cells from the solid phase are known in the art and encompass, for example, the use of enzymes. Such enzymes are preferably non-toxic and non-injurious to the cells and are preferably directed to cleaving the cell-surface binding partner. Alternatively, mixtures of cells suspected of containing polypeptide-expressing cells of the invention first can be incubated with a biotinylated polypeptide-binding polypeptide of the invention. The resulting mixture then is passed through a column packed with avidin-coated beads, whereby the high affinity of biotin for avidin provides the binding of the polypeptide-binding cells to the beads. Use of avidin-coated beads is known in the art. See Berenson, et al. J. Cell. Biochem., 10D:239 (1986). Wash of unbound material and the release of the bound cells is performed using conventional methods.

The polypeptide can also be produced by known conventional chemical synthesis. The synthetically-constructed polypeptide sequences, by virtue of sharing primary, secondary or tertiary structural and/or conformational characteristics with HPR1 and/or HPR2 polypeptides can possess biological properties in common therewith, including HPR1 and/or HPR2 polypeptide activity. Thus, they can be employed as biologically active or immunological substitutes for natural, purified polypeptides in screening of therapeutic compounds and in immunological processes for the development of antibodies.

The desired degree of purity depends on the intended use of the polypeptide. A relatively high degree of purity is desired when the polypeptide is to be administered *in vivo*, for example. In such a case, the polypeptides are purified such that no polypeptide bands corresponding to other polypeptides are detectable upon analysis by SDS-polyacrylamide gel electrophoresis (SDS-PAGE). It will be recognized by one skilled in the pertinent field that multiple bands corresponding to the polypeptide can be visualized by SDS-PAGE, due to differential glycosylation, differential post-translational processing, and the like. Most preferably, the polypeptide of the invention is purified to substantial homogeneity, as indicated by a single polypeptide band upon analysis by SDS-PAGE. The polypeptide band can be visualized by silver staining, Coomassie blue staining, or (if the polypeptide is radiolabeled) by autoradiography.

### Antagonists and Agonists of HPR1 and/or HPR2 Polypeptides

Any method which neutralizes HPR1 and/or HPR2 polypeptides or inhibits expression of the HPR1 and/or HPR2 genes (either transcription or translation) can be used to reduce the biological activities of HPR1 and/or HPR2 polypeptides. In particular embodiments, antagonists inhibit the binding of at least one HPR1 polypeptide and/or at least one HPR2 polypeptide to cells, thereby inhibiting biological activities induced by the binding of those HPR1 or HPR2 polypeptides to the

5 cells. In certain other embodiments of the invention, antagonists can be designed to reduce the level of endogenous HPR1 and/or HPR2 gene expression, e.g., using well-known antisense or ribozyme approaches to inhibit or prevent translation of HPR1 and/or HPR2 mRNA transcripts; triple helix approaches to inhibit transcription of HPR1 and/or HPR2 genes; or targeted homologous recombination to inactivate or "knock out" the HPR1 gene(s), the HPR2 gene(s), or their endogenous promoters or enhancer elements. Such antisense, ribozyme, and triple helix antagonists can be designed to reduce or inhibit either unimpaired, or if appropriate, mutant HPR1 and/or HPR2 gene activity. Techniques for the production and use of such molecules are well known to those of skill in the art.

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Antisense RNA and DNA molecules act to directly block the translation of mRNA by hybridizing to targeted mRNA and preventing polypeptide translation. Antisense approaches involve the design of oligonucleotides (either DNA or RNA) that are complementary to an HPR1 and/or to an HPR2 mRNA. The antisense oligonucleotides will bind to the complementary target gene mRNA transcripts and prevent translation. Absolute complementarity, although preferred, is not required. A sequence "complementary" to a portion of a nucleic acid, as referred to herein, means a sequence having sufficient complementarity to be able to hybridize with the nucleic acid, forming a stable duplex (or triplex, as appropriate). In the case of double-stranded antisense nucleic acids, a single strand of the duplex DNA can thus be tested, or triplex formation can be assayed. The ability to hybridize will depend on both the degree of complementarity and the length of the antisense nucleic acid. Preferred oligonucleotides are complementary to the 5' end of the message, e.g., the 5' untranslated sequence up to and including the AUG initiation codon. However, oligonucleotides complementary to the 5'- or 3'non-translated, non-coding regions of the HPR1 or HPR2 gene transcript(s) could be used in an antisense approach to inhibit translation of endogenous HPR1 and/or HPR2 mRNA. Antisense nucleic acids should be at least six nucleotides in length, and are preferably oligonucleotides ranging from 6 to about 50 nucleotides in length. In specific aspects the oligonucleotide is at least 10 nucleotides, at least 17 nucleotides, at least 25 nucleotides or at least 50 nucleotides. The oligonucleotides can be DNA or RNA or chimeric mixtures or derivatives or modified versions thereof, single-stranded or doublestranded. Chimeric oligonucleotides, oligonucleosides, or mixed oligonucleotides/oligonucleosides of the invention can be of several different types. These include a first type wherein the "gap" segment of nucleotides is positioned between 5' and 3' "wing" segments of linked nucleosides and a second "open end" type wherein the "gap" segment is located at either the 3' or the 5' terminus of the oligomeric compound (see, e.g., U.S. Pat. No. 5,985,664). Oligonucleotides of the first type are also known in the art as "gapmers" or gapped oligonucleotides. Oligonucleotides of the second type are also known in the art as "hemimers" or "wingmers". The oligonucleotide can be modified at the base moiety, sugar moiety, or phosphate backbone, for example, to improve stability of the molecule, hybridization, etc. The oligonucleotide can include other appended groups such as peptides (e.g., for targeting host cell receptors in vivo), or agents facilitating transport across the cell membrane (see, e.g., Letsinger et al., 1989, Proc Natl Acad Sci USA 86:6553-6556; Lemaitre et al., 1987, Proc Natl Acad Sci 84:648-652;

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PCT Publication No. WO88/09810), or hybridization-triggered cleavage agents or intercalating agents. (See, e.g., Zon, 1988, Pharm. Res. 5:539-549). The antisense molecules should be delivered to cells which express the HPR1 and/or HPR2 transcript in vivo. A number of methods have been developed for delivering antisense DNA or RNA to cells; e.g., antisense molecules can be injected directly into the tissue or cell derivation site, or modified antisense molecules, designed to target the desired cells (e.g., antisense linked to peptides or antibodies that specifically bind receptors or antigens expressed on the target cell surface) can be administered systemically. However, it is often difficult to achieve intracellular concentrations of the antisense sufficient to suppress translation of endogenous mRNAs. Therefore a preferred approach utilizes a recombinant DNA construct in which the antisense oligonucleotide is placed under the control of a strong pol III or pol II promoter. The use of such a construct to transfect target cells in the patient will result in the transcription of sufficient amounts of single stranded RNAs that will form complementary base pairs with the endogenous HPR1 and/or HPR2 gene transcripts and thereby prevent translation of the HPR1 and/or HPR2 mRNA. For example, a vector can be introduced in vivo such that it is taken up by a cell and directs the transcription of an antisense RNA. Such a vector can remain episomal or become chromosomally integrated, as long as it can be transcribed to produce the desired antisense RNA. Such vectors can be constructed by recombinant DNA technology methods standard in the art. Vectors can be plasmid, viral, or others known in the art, used for replication and expression in mammalian cells.

Ribozyme molecules designed to catalytically cleave HPR1 and/or HPR2 mRNA transcripts can also be used to prevent translation of HPR1 and/or HPR2 mRNA and expression of HPR1 and/or HPR2 polypeptides. (See, e.g., PCT International Publication WO90/11364 and US Patent No. 5,824,519). The ribozymes that can be used in the present invention include hammerhead ribozymes (Haseloff and Gerlach, 1988, Nature, 334:585-591), RNA endoribonucleases (hereinafter "Cech-type ribozymes") such as the one which occurs naturally in Tetrahymena Thermophila (known as the IVS, or L-19 IVS RNA) and which has been extensively described by Thomas Cech and collaborators (International Patent Application No. WO 88/04300; Been and Cech, 1986, Cell, 47:207-216). As in the antisense approach, the ribozymes can be composed of modified oligonucleotides (e.g. for improved stability, targeting, etc.) and should be delivered to cells which express HPR1 and/or HPR2 polypeptides in vivo. A preferred method of delivery involves using a DNA construct "encoding" the ribozyme under the control of a strong constitutive pol III or pol II promoter, so that transfected cells will produce sufficient quantities of the ribozyme to destroy endogenous HPR1 and/or HPR2 messages and inhibit translation. Because ribozymes, unlike antisense molecules, are catalytic, a lower intracellular concentration is required for efficiency.

Alternatively, endogenous HPR1 and/or HPR2 gene expression can be reduced by targeting deoxyribonucleotide sequences complementary to the regulatory region of the target gene (i.e., the target gene promoter and/or enhancers) to form triple helical structures that prevent transcription of the target HPR1 and/or HPR2 gene. (See generally, Helene, 1991, Anticancer Drug Des., 6(6), 569-584; Helene, et al., 1992, Ann. N.Y. Acad. Sci., 660, 27-36; and Maher, 1992, Bioassays 14(12), 807-815).

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Anti-sense RNA and DNA, ribozyme, and triple helix molecules of the invention can be prepared by any method known in the art for the synthesis of DNA and RNA molecules. These include techniques for chemically synthesizing oligodeoxyribonucleotides and oligoribonucleotides well known in the art such as for example solid phase phosphoramidite chemical synthesis. Oligonucleotides can be synthesized by standard methods known in the art, e.g. by use of an automated DNA synthesizer (such as are commercially available from Biosearch, Applied Biosystems, etc.). As examples, phosphorothicate oligonucleotides can be synthesized by the method of Stein et al., 1988, Nucl. Acids Res. 16:3209. Methylphosphonate oligonucleotides can be prepared by use of controlled pore glass polymer supports (Sarin et al., 1988, Proc. Natl. Acad. Sci. U.S.A. 85:7448-7451). Alternatively, RNA molecules can be generated by in vitro and in vivo transcription of DNA sequences encoding the antisense RNA molecule. Such DNA sequences can be incorporated into a wide variety of vectors that incorporate suitable RNA polymerase promoters such as the T7 or SP6 polymerase promoters. Alternatively, antisense cDNA constructs that synthesize antisense RNA constitutively or inducibly, depending on the promoter used, can be introduced stably into cell lines.

Endogenous target gene expression can also be reduced by inactivating or "knocking out" the target gene or its promoter using targeted homologous recombination (e.g., see Smithies, et al., 1985, Nature 317, 230-234; Thomas and Capecchi, 1987, Cell 51, 503-512; Thompson, et al., 1989, Cell 5, 313-321). For example, a mutant, non-functional target gene (or a completely unrelated DNA sequence) flanked by DNA homologous to the endogenous target gene (either the coding regions or regulatory regions of the target gene) can be used, with or without a selectable marker and/or a negative selectable marker, to transfect cells that express the target gene in vivo. Insertion of the DNA construct, via targeted homologous recombination, results in inactivation of the target gene. Such approaches are particularly suited in the agricultural field where modifications to ES (embryonic stem) cells can be used to generate animal offspring with an inactive target gene (e.g., see Thomas and Capecchi, 1987 and Thompson, 1989, supra), or in model organisms such as Caenorhabditis elegans where the "RNA interference" ("RNAi") technique (Grishok, Tabara, and Mello, 2000, Genetic requirements for inheritance of RNAi in C. elegans, Science 287 (5462): 2494-2497), or the introduction of transgenes (Dernburg et al., 2000, Transgene-mediated cosuppression in the C. elegans germ line, Genes Dev. 14 (13): 1578-1583) are used to inhibit the expression of specific target genes. However this approach can be adapted for use in humans provided the recombinant DNA constructs are directly administered or targeted to the required site in vivo using appropriate vectors such as viral vectors.

Organisms that have enhanced, reduced, or modified expression of the gene(s) corresponding to the nucleic acid sequences disclosed herein are provided. The desired change in gene expression can be achieved through the use of antisense nucleic acids or ribozymes that bind and/or cleave the mRNA transcribed from the gene (Albert and Morris, 1994, Trends Pharmacol. Sci. 15(7): 250-254; Lavarosky et al., 1997, Biochem. Mol. Med. 62(1): 11-22; and Hampel, 1998, Prog. Nucleic Acid Res. Mol. Biol. 58: 1-39). Transgenic animals that have multiple copies of the gene(s) corresponding to the nucleic

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acid sequences disclosed herein, preferably produced by transformation of cells with genetic constructs that are stably maintained within the transformed cells and their progeny, are provided. Transgenic animals that have modified genetic control regions that increase or reduce gene expression levels, or that change temporal or spatial patterns of gene expression, are also provided (see European Patent No. 0 649 464 B1). In addition, organisms are provided in which the gene(s) corresponding to the nucleic acid sequences disclosed herein have been partially or completely inactivated, through insertion of extraneous sequences into the corresponding gene(s) or through deletion of all or part of the corresponding gene(s). Partial or complete gene inactivation can be accomplished through insertion, preferably followed by imprecise excision, of transposable elements (Plasterk, 1992, Bioessays 14(9): 629-633; Zwaal et al., 1993, Proc Natl Acad Sci USA 90(16): 7431-7435; Clark et al., 1994, Proc Natl Acad Sci USA 91(2): 719-722), or through homologous recombination, preferably detected by positive/negative genetic selection strategies (Mansour et al., 1988, Nature 336: 348-352; U.S. Pat. Nos. 5,464,764; 5,487,992; 5,627,059; 5,631,153; 5,614,396; 5,616,491; and 5,679,523). These organisms with altered gene expression are preferably eukaryotes and more preferably are mammals. Such organisms are useful for the development of non-human models for the study of disorders involving the corresponding gene(s), and for the development of assay systems for the identification of molecules that interact with the polypeptide product(s) of the corresponding gene(s).

Also encompassed within the invention are HPR1 and HPR2 polypeptide variants with partner binding sites that have been altered in conformation so that (1) the HPR1 or HPR2 variant will still bind to its partner(s), but a specified small molecule will fit into the altered binding site and block that interaction, or (2) the HPR1 or HPR2 variant will no longer bind to its partner(s) unless a specified small molecule is present (see for example Bishop et al., 2000, Nature 407: 395-401). Nucleic acids encoding such altered HPR1 or HPR2 polypeptides can be introduced into organisms according to methods described herein, and can replace the endogenous nucleic acid sequences encoding the corresponding HPR1 or HPR2 polypeptide. Such methods allow for the interaction of a particular HPR1 or HPR2 polypeptide with its binding partners to be regulated by administration of a small molecule compound to an organism, either systemically or in a localized manner.

The HPR1 and HPR2 polypeptides themselves can also be employed in inhibiting a biological activity of HPR1 and /or of HPR2 in *in vitro* or *in vivo* procedures. Encompassed within the invention are cytokine receptor domains of HPR1 and HPR2 polypeptides that act as "dominant negative" inhibitors of native HPR1 and/or HPR2 polypeptide function when expressed as fragments or as components of fusion polypeptides. For example, a purified polypeptide domain of the present invention can be used to inhibit binding of HPR1 or HPR2 polypeptides to endogenous binding partners. Such use effectively would block HPR1 and/or HPR2 polypeptide interactions and inhibit HPR1 and/or HPR2 polypeptide activities. In still another aspect of the invention, a soluble form of an HPR1 and/or HPR2 binding partner is used to bind to an endogenous HPR1 and/or HPR2 polypeptide, and competitively inhibit activation of that endogenous HPR1 and/or HPR2 polypeptide. Furthermore, antibodies which bind to HPR1 and/or HPR2 polypeptides often inhibit HPR1 and/or HPR2

polypeptide activity and act as antagonists. For example, antibodies that specifically recognize one or more epitopes of HPR1 and/or HPR2 polypeptides, or epitopes of conserved variants of HPR1 and/or HPR2 polypeptides, or peptide fragments of an HPR1 and/or HPR2 polypeptide can be used in the invention to inhibit HPR1 and/or HPR2 polypeptide activity. Such antibodies include but are not limited to polyclonal antibodies, monoclonal antibodies (mAbs), humanized or chimeric antibodies, single chain antibodies, Fab fragments, F(ab')2 fragments, fragments produced by a Fab expression library, anti-idiotypic (anti-Id) antibodies, and epitope-binding fragments of any of the above. Alternatively, purified and modified HPR1 and/or HPR2 polypeptides of the present invention can be administered to modulate interactions between HPR1 and/or HPR2 polypeptides and HPR1 and/or HPR2 binding partners that are not membrane-bound. Such an approach will allow an alternative method for the modification of HPR1- and/or HPR2-influenced bioactivity.

In an alternative aspect, the invention further encompasses the use of agonists of HPR1 and/or HPR2 polypeptide activity to treat or ameliorate the symptoms of a disease for which increased HPR1 and/or HPR2 polypeptide activity is beneficial. Such diseases include but are not limited to pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, osteoporosis resulting from a lack of bone-forming cells, obesity, deficient mammary development, and infertility. In a preferred aspect, the invention entails administering compositions comprising an HPR1 or HPR2 nucleic acid or an HPR1 or HPR2 polypeptide to cells in vitro, to cells ex vivo, to cells in vivo, and/or to a multicellular organism such as a vertebrate or mammal. Preferred therapeutic forms of HPR1 and HPR2 are soluble forms, as described above. In still another aspect of the invention, the compositions comprise administering an HPR1-encoding nucleic acid or an HPR2-encoding nucleic acid for expression of an HPR1 or HPR2 polypeptide in a host organism for treatment of disease. Particularly preferred in this regard is expression in a human patient for treatment of a dysfunction associated with aberrant (e.g., decreased) endogenous activity of an HPR1 or HPR2 polypeptide. Furthermore, the invention encompasses the administration to cells and/or organisms of compounds found to increase the endogenous activity of HPR1 and/or HPR2 polypeptides. One example of compounds that increase HPR1 and/or HPR2 polypeptide activity are agonistic antibodies, preferably monoclonal antibodies, that bind to HPR1 and/or HPR2 polypeptides or binding partners, which may increase HPR1 and/or HPR2 polypeptide activity by causing constitutive intracellular signaling (or "ligand mimicking"), or by preventing the binding of a native inhibitor of HPR1 and/or HPR2 polypeptide activity.

Antibodies to HPR1 and/or HPR2 Polypeptides

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Antibodies that are immunoreactive with the polypeptides of the invention are provided herein. Such antibodies specifically bind to the polypeptides via the antigen-binding sites of the antibody (as opposed to non-specific binding). In the present invention, specifically binding antibodies are those that will specifically recognize and bind with HPR1 and/or HPR2 polypeptides, homologues, and variants, but not with other molecules. In one preferred embodiment, the antibodies are specific for

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the polypeptides of the present invention and do not cross-react with other polypeptides. In this manner, the HPR1 and HPR2 polypeptides, fragments, variants, fusion polypeptides, etc., as set forth above can be employed as "immunogens" in producing antibodies immunoreactive therewith.

More specifically, the polypeptides, fragment, variants, fusion polypeptides, etc. contain antigenic determinants or epitopes that elicit the formation of antibodies. These antigenic determinants or epitopes can be either linear or conformational (discontinuous). Linear epitopes are composed of a single section of amino acids of the polypeptide, while conformational or discontinuous epitopes are composed of amino acids sections from different regions of the polypeptide chain that are brought into close proximity upon polypeptide folding (Janeway and Travers, Immuno Biology 3:9 (Garland Publishing Inc., 2nd ed. 1996)). Because folded polypeptides have complex surfaces, the number of epitopes available is quite numerous; however, due to the conformation of the polypeptide and steric hinderances, the number of antibodies that actually bind to the epitopes is less than the number of available epitopes (Janeway and Travers, Immuno Biology 2:14 (Garland Publishing Inc., 2nd ed. 1996)). Epitopes can be identified by any of the methods known in the art. Thus, one aspect of the present invention relates to the antigenic epitopes of the polypeptides of the invention. Such epitopes are useful for raising antibodies, in particular monoclonal antibodies, as described in more detail below. Additionally, epitopes from the polypeptides of the invention can be used as research reagents, in assays, and to purify specific binding antibodies from substances such as polyclonal sera or supernatants from cultured hybridomas. Such epitopes or variants thereof can be produced using techniques well known in the art such as solid-phase synthesis, chemical or enzymatic cleavage of a polypeptide, or using recombinant DNA technology.

As to the antibodies that can be elicited by the epitopes of the polypeptides of the invention, whether the epitopes have been isolated or remain part of the polypeptides, both polyclonal and monoclonal antibodies can be prepared by conventional techniques. See, for example, Monoclonal Antibodies, Hybridomas: A New Dimension in Biological Analyses, Kennet et al. (eds.), Plenum Press, New York (1980); and Antibodies: A Laboratory Manual, Harlow and Land (eds.), Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, (1988); Kohler and Milstein, (U.S. Pat. No. 4,376,110); the human B-cell hybridoma technique (Kosbor et al., 1984, J Immunol 133: 3001-3005; Cole et al., 1983, Proc Natl Acad Sci USA 80:2026-2030); and the EBV-hybridoma technique (Cole et al., 1985, Monoclonal Antibodies And Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). Hybridoma cell lines that produce monoclonal antibodies specific for the polypeptides of the invention are also contemplated herein. Such hybridomas can be produced and identified by conventional techniques. The hybridoma producing the mAb of this invention can be cultivated in vitro or in vivo. Production of high titers of mAbs in vivo makes this the presently preferred method of production. One method for producing such a hybridoma cell line comprises immunizing an animal with a polypeptide; harvesting spleen cells from the immunized animal; fusing said spleen cells to a myeloma cell line, thereby generating hybridoma cells; and identifying a hybridoma cell line that produces a monoclonal antibody that binds the polypeptide. For the production of antibodies, various host animals can be immunized by

injection with one or more of the following: an HPR1 or HPR2 polypeptide, a fragment of an HPR1 or HPR2 polypeptide, a functional equivalent of an HPR1 or HPR2 polypeptide, or a mutant form of an HPR1 or HPR2 polypeptide. Such host animals can include but are not limited to rabbits, mice, and rats. Various adjuvants can be used to increase the immunologic response, depending on the host species, including but not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanin, dinitrophenol, and potentially useful human adjutants such as BCG (bacille Calmette-Guerin) and Corynebacterium parvum. The monoclonal antibodies can be recovered by conventional techniques. Such monoclonal antibodies can be of any immunoglobulin class including IgG, IgM, IgE, IgA, IgD and any subclass thereof.

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In addition, techniques developed for the production of "chimeric antibodies" (Takeda et al., 1985, Nature, 314: 452-454; Morrison et al., 1984, Proc Natl Acad Sci USA 81: 6851-6855; Boulianne et al., 1984, Nature 312: 643-646; Neuberger et al., 1985, Nature 314: 268-270) by splicing the genes from a mouse antibody molecule of appropriate antigen specificity together with genes from a human antibody molecule of appropriate biological activity can be used. A chimeric antibody is a molecule in which different portions are derived from different animal species, such as those having a variable region derived from a porcine mAb and a human immunoglobulin constant region. The monoclonal antibodies of the present invention also include humanized versions of murine monoclonal antibodies. Such humanized antibodies can be prepared by known techniques and offer the advantage of reduced immunogenicity when the antibodies are administered to humans. In one embodiment, a humanized monoclonal antibody comprises the variable region of a murine antibody (or just the antigen binding site thereof) and a constant region derived from a human antibody. Alternatively, a humanized antibody fragment can comprise the antigen binding site of a murine monoclonal antibody and a variable region fragment (lacking the antigen-binding site) derived from a human antibody. Procedures for the production of chimeric and further engineered monoclonal antibodies include those described in Riechmann et al. (Nature 332:323, 1988), Liu et al. (PNAS 84:3439, 1987), Larrick et al. (Bio/Technology 7:934, 1989), and Winter and Harris (TIPS 14:139, Can, 1993). Useful techniques for humanizing antibodies are also discussed in U.S. Patent 6,054,297. Procedures to generate antibodies transgenically can be found in GB 2,272,440, US Patent Nos. 5,569,825 and 5,545,806, and related patents. Preferably, for use in humans, the antibodies are human or humanized; techniques for creating such human or humanized antibodies are also well known and are commercially available from, for example, Medarex Inc. (Princeton, NJ) and Abgenix Inc. (Fremont, CA). In another preferred embodiment, fully human antibodies for use in humans are produced by screening a phage display library of human antibody variable domains (Vaughan et al., 1998, Nat Biotechnol. 16(6): 535-539; and U.S. Patent No. 5,969,108).

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Antigen-binding antibody fragments which recognize specific epitopes can be generated by known techniques. For example, such fragments include but are not limited to: the F(ab')2 fragments which can be produced by pepsin digestion of the antibody molecule and the Fab fragments which can

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be generated by reducing the disulfide bridges of the (ab')2 fragments. Alternatively, Fab expression libraries can be constructed (Huse *et al.*, 1989, Science, 246:1275-1281) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity. Techniques described for the production of single chain antibodies (U.S. Pat. No. 4,946,778; Bird, 1988, Science 242:423-426; Huston *et al.*, 1988, Proc. Natl. Acad. Sci. USA 85:5879-5883; and Ward *et al.*, 1989, Nature 334:544-546) can also be adapted to produce single chain antibodies against HPR1 and/or HPR2 gene products. Single chain antibodies are formed by linking the heavy and light chain fragments of the Fv region via an amino acid bridge, resulting in a single chain polypeptide. Such single chain antibodies can also be useful intracellularly (i.e., as 'intrabodies), for example as described by Marasco *et al.* (*J. Immunol. Methods* 231:223-238, 1999) for genetic therapy in HIV infection. In addition, antibodies to the HPR1 and/or HPR2 polypeptide can, in turn, be utilized to generate anti-idiotype antibodies that "mimic" the HPR1 and/or HPR2 polypeptide and that may bind to the binding partner(s) of HPR1 and/or HPR2 polypeptides, using techniques well known to those skilled in the art. (See, *e.g.*, Greenspan & Bona, 1993, FASEB J 7(5):437-444; and Nissinoff, 1991, J. Immunol. 147(8):2429-2438).

Antibodies that are immunoreactive with the polypeptides of the invention include bispecific antibodies (i.e., antibodies that are immunoreactive with the polypeptides of the invention via a first antigen binding domain, and also immunoreactive with a different polypeptide via a second antigen binding domain). A variety of bispecific antibodies have been prepared, and found useful both in vitro and in vivo (see, for example, U.S. Patent 5,807,706; and Cao and Suresh, 1998, Bioconjugate Chem 9: 635-644). Numerous methods of preparing bispecific antibodies are known in the art, including the use of hybrid-hybridomas such as quadromas, which are formed by fusing two differed hybridomas, and triomas, which are formed by fusing a hybridoma with a lymphocyte (Milstein and Cuello, 1983, Nature 305: 537-540; U.S. Patent 4,474,893; and U.S. Patent 6,106,833). U.S. Patent 6,060,285 discloses a process for the production of bispecific antibodies in which at least the genes for the light chain and the variable portion of the heavy chain of an antibody having a first specificity are transfected into a hybridoma cell secreting an antibody having a second specificity. Chemical coupling of antibody fragments has also been used to prepare antigen-binding molecules having specificity for two different antigens (Brennan et al., 1985, Science 229: 81-83; Glennie et al., J. Immunol., 1987, 139:2367-2375; and U.S. Patent 6,010,902). Bispecific antibodies can also be produced via recombinant means, for example, by using the leucine zipper moieties from the Fos and Jun proteins (which preferentially form heterodimers) as described by Kostelny et al. (J. Immnol. 148:1547-4553; 1992). U.S. Patent 5,582,996 discloses the use of complementary interactive domains (such as leucine zipper moieties or other lock and key interactive domain structures) to facilitate heterodimer formation in the production of bispecific antibodies. Tetravalent, bispecific molecules can be prepared by fusion of DNA encoding the heavy chain of an F(ab')2 fragment of an antibody with either DNA encoding the heavy chain of a second F(ab')2 molecule (in which the CH1 domain is replaced by a CH3 domain), or with DNA encoding a single chain FV fragment of an antibody, as described in U.S. Patent 5,959,083. Expression of the resultant fusion genes in mammalian cells, together with the genes for the

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corresponding light chains, yields tetravalent bispecific molecules having specificity for selected antigens. Bispecific antibodies can also be produced as described in U.S. Patent 5,807,706. Generally, the method involves introducing a protuberance (constructed by replacing small amino acid side chains with larger side chains) at the interface of a first polypeptide and a corresponding cavity (prepared by replacing large amino acid side chains with smaller ones) in the interface of a second polypeptide. Moreover, single-chain variable fragments (sFvs) have been prepared by covalently joining two variable domains; the resulting antibody fragments can form dimers or trimers, depending on the length of a flexible linker between the two variable domains (Kortt *et al.*, 1997, *Protein Engineering* 10:423-433).

Screening procedures by which such antibodies can be identified are well known, and can involve immunoaffinity chromatography, for example. Antibodies can be screened for agonistic (i.e., ligand-mimicking) properties. Such antibodies, upon binding to cell surface HPR1 and/or HPR2, induce biological effects (e.g., transduction of biological signals) similar to the biological effects induced when the HPR1 and/or HPR2 binding partner binds to cell surface HPR1 and/or HPR2. Agonistic antibodies can be used to induce HPR1- and/or HPR2-mediated intracellular signaling or cell proliferation. Bispecific antibodies can be identified by screening with two separate assays, or with an assay wherein the bispecific antibody serves as a bridge between the first antigen and the second antigen (the latter is coupled to a detectable moiety). Bispecific antibodies that bind HPR1 and/or HPR2 polypeptides of the invention via a first antigen binding domain will be useful in diagnostic applications and in treating cell proliferation, differentiation, or activation diseases or conditions. Examples of polypeptides (or other antigens) that the inventive bispecific antibodies bind via a second antigen binding domain include: four alpha helix bundle cytokines such as IL-6, OSM, LIF, CNTF, CLC, IL-12p35, and IL-23p19; soluble hematopoietin receptors such as EBI-3, soluble IL-6R alpha, cytokine-like factor-1 (CLF), IL-12p40, or a soluble form of HPR1 and/or HPR2; and soluble hematopoietin receptors such as EBI-3 etc. in conjunction with a four alpha helix bundle cytokine.

Those antibodies that can block binding of the HPR1 and/or HPR2 polypeptides of the invention to binding partners for HPR1 and/or HPR2 can be used to inhibit HPR1- and/or HPR2-mediated intracellular signaling or cell proliferation that results from such binding. Such blocking antibodies can be identified using any suitable assay procedure, such as by testing antibodies for the ability to inhibit binding of HPR1 and/or HPR2 to certain cells expressing an HPR1 and/or HPR2 binding partner. Alternatively, blocking antibodies can be identified in assays for the ability to inhibit a biological effect that results from binding of soluble HPR1 and/or HPR2 to target cells. Antibodies can be assayed for the ability to inhibit HPR1 and/or HPR2 binding partner-mediated cell stimulatory pathways, for example. Such an antibody can be employed in an *in vitro* procedure, or administered *in vivo* to inhibit a biological activity mediated by the entity that generated the antibody. Disorders caused or exacerbated (directly or indirectly) by the interaction of HPR1 and/or HPR2 with cell surface binding partner receptor thus can be treated. A therapeutic method involves *in vivo* administration of a blocking antibody to a mammal in an amount effective in inhibiting HPR1 and/or HPR2 binding

partner-mediated biological activity. Monoclonal antibodies are generally preferred for use in such therapeutic methods. In one embodiment, an antigen-binding antibody fragment is employed. Compositions comprising an antibody that is directed against HPR1 and/or HPR2, and a physiologically acceptable diluent, excipient, or carrier, are provided herein. Suitable components of such compositions are as described below for compositions containing HPR1 and/or HPR2 polypeptides.

Also provided herein are conjugates comprising a detectable (e.g., diagnostic) or therapeutic agent, attached to the antibody. Examples of such agents are presented above. The conjugates find use in *in vitro* or *in vivo* procedures. The antibodies of the invention can also be used in assays to detect the presence of the polypeptides or fragments of the invention, either *in vitro* or *in vivo*. The antibodies also can be employed in purifying polypeptides or fragments of the invention by immunoaffinity chromatography.

#### Rational Design of Compounds that Interact with HPR1 and/or HPR2 Polypeptides

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The goal of rational drug design is to produce structural analogs of biologically active polypeptides of interest or of small molecules with which they interact, e.g., inhibitors, agonists, antagonists, etc. Any of these examples can be used to fashion drugs which are more active or stable forms of the polypeptide or which enhance or interfere with the function of a polypeptide in vivo (Hodgson J (1991) Biotechnology 9:19-21). In one approach, the three-dimensional structure of a polypeptide of interest, or of a polypeptide-inhibitor complex, is determined by x-ray crystallography, by nuclear magnetic resonance, or by computer homology modeling or, most typically, by a combination of these approaches. Both the shape and charges of the polypeptide must be ascertained to elucidate the structure and to determine active site(s) of the molecule. Less often, useful information regarding the structure of a polypeptide may be gained by modeling based on the structure of homologous polypeptides. In both cases, relevant structural information is used to design analogous HPR1- and/or HPR2-like molecules, to identify efficient inhibitors, or to identify small molecules that bind HPR1 and/or HPR2 polypeptides. Useful examples of rational drug design include molecules which have improved activity or stability as shown by Braxton S and Wells JA (1992 Biochemistry 31:7796-7801) or which act as inhibitors, agonists, or antagonists of native peptides as shown by Athauda SB et al (1993 J Biochem 113:742-746). The use of HPR1 and/or HPR2 polypeptide structural information in molecular modeling software systems to assist in inhibitor design and in studying inhibitor-HPR1 polypeptide and/or inhibitor-HPR2 polypeptide interaction is also encompassed by the invention. A particular method of the invention comprises analyzing the threedimensional structure of HPR1 and/or HPR2 polypeptides for likely binding sites of substrates, synthesizing a new molecule that incorporates a predictive reactive site, and assaying the new molecule as described further herein.

It is also possible to isolate a target-specific antibody, selected by functional assay, as described further herein, and then to solve its crystal structure. This approach, in principle, yields a pharmacore upon which subsequent drug design can be based. It is possible to bypass polypeptide

crystallography altogether by generating anti-idiotypic antibodies (anti-ids) to a functional, pharmacologically active antibody. As a mirror image of a mirror image, the binding site of the anti-ids would be expected to be an analog of the original receptor. The anti-id could then be used to identify and isolate peptides from banks of chemically or biologically produced peptides. The isolated peptides would then act as the pharmacore.

## 10 Assays of HPR1 and HPR2 Polypeptide Activities

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The purified HPR1 and HPR2 polypeptides of the invention (including polypeptides, polypeptides, fragments, variants, oligomers, and other forms) are useful in a variety of assays. For example, the HPR1 and HPR2 molecules of the present invention can be used to identify binding partners of HPR1 and/or HPR2 polypeptides, which can also be used to modulate intracellular signaling, cell proliferation, or immune cell activity. Alternatively, they can be used to identify non-binding-partner molecules or substances that modulate intracellular signaling, cell proliferation, or immune cell activity.

Assays to Identify Binding Partners. HPR1 and HPR2 polypeptides and fragments thereof can be used to identify binding partners. For example, they can be tested for the ability to bind a candidate binding partner in any suitable assay, such as a conventional binding assay. To illustrate, the HPR1 or HPR2 polypeptide can be labeled with a detectable reagent (e.g., a radionuclide, chromophore, enzyme that catalyzes a colorimetric or fluorometric reaction, and the like). The labeled polypeptide is contacted with cells expressing the candidate binding partner. The cells then are washed to remove unbound labeled polypeptide, and the presence of cell-bound label is determined by a suitable technique, chosen according to the nature of the label.

One example of a binding assay procedure is as follows. A recombinant expression vector containing the candidate binding partner cDNA is constructed. CV1-EBNA-1 cells in 10 cm<sup>2</sup> dishes are transfected with this recombinant expression vector. CV-1/EBNA-1 cells (ATCC CRL 10478) constitutively express EBV nuclear antigen-1 driven from the CMV Immediate-early enhancer/promoter. CV1-EBNA-1 was derived from the African Green Monkey kidney cell line CV-1 (ATCC CCL 70), as described by McMahan et al., (EMBO J. 10:2821, 1991). The transfected cells are cultured for 24 hours, and the cells in each dish then are split into a 24-well plate. After culturing an additional 48 hours, the transfected cells (about 4 x 10<sup>4</sup> cells/well) are washed with BM-NFDM, which is binding medium (RPMI 1640 containing 25 mg/ml bovine serum albumin, 2 mg/ml sodium azide, 20 mM Hepes pH 7.2) to which 50 mg/ml nonfat dry milk has been added. The cells then are incubated for 1 hour at 37°C with various concentrations of, for example, a soluble polypeptide/Fc fusion polypeptide made as set forth above. Cells then are washed and incubated with a constant saturating concentration of a <sup>125</sup>I-mouse anti-human IgG in binding medium, with gentle agitation for 1 hour at 37°C. After extensive washing, cells are released via trypsinization. The mouse anti-human IgG employed above is directed against the Fc region of human IgG and can be obtained from Jackson Immunoresearch Laboratories, Inc., West Grove, PA. The antibody is radioiodinated using the standard chloramine-T method. The antibody will bind to the Fc portion of any polypeptide/Fc

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polypeptide that has bound to the cells. In all assays, non-specific binding of <sup>125</sup>I-antibody is assayed in the absence of the Fc fusion polypeptide/Fc, as well as in the presence of the Fc fusion polypeptide and a 200-fold molar excess of unlabeled mouse anti-human IgG antibody. Cell-bound 125I-antibody is quantified on a Packard Autogamma counter. Affinity calculations (Scatchard, Ann. N.Y. Acad. Sci. 51:660, 1949) are generated on RS/1 (BBN Software, Boston, MA) run on a Microvax computer. Binding can also be detected using methods that are well suited for high-throughput screening procedures, such as scintillation proximity assays (Udenfriend et al., 1985, Proc Natl Acad Sci USA 82: 8672-8676), homogeneous time-resolved fluorescence methods (Park et al., 1999, Anal Biochem 269: 94-104), fluorescence resonance energy transfer (FRET) methods (Clegg RM, 1995, Curr Opin Biotechnol 6: 103-110), or methods that measure any changes in surface plasmon resonance when a bound polypeptide is exposed to a potential binding partner, using for example a biosensor such as that supplied by Biacore AB (Uppsala, Sweden). Compounds that can be assayed for binding to HPR1 and/or HPR2 polypeptides include but are not limited to small organic molecules, such as those that are commercially available - often as part of large combinatorial chemistry compound 'libraries' - from companies such as Sigma-Aldrich (St. Louis, MO), Arqule (Woburn, MA), Enzymed (Iowa City, IA), Maybridge Chemical Co.(Trevillett, Cornwall, UK), MDS Panlabs (Bothell, WA), Pharmacopeia (Princeton, NJ), and Trega (San Diego, CA). Preferred small organic molecules for screening using these assays are usually less than 10K molecular weight and can possess a number of physicochemical and pharmacological properties which enhance cell penetration, resist degradation, and/or prolong their physiological half-lives (Gibbs, J., 1994, Pharmaceutical Research in Molecular Oncology, Cell 79(2): 193-198). Compounds including natural products, inorganic chemicals, and biologically active materials such as proteins and toxins can also be assayed using these methods for the ability to bind to HPR1 and/or HPR2 polypeptides.

Yeast Two-Hybrid or "Interaction Trap" Assays. Because HPR1 and HPR2 polypeptides bind or potentially bind to another polypeptide (such as, for example, in a receptor-ligand interaction), the nucleic acid encoding the HPR1 or HPR2 polypeptide can also be used in interaction trap assays (such as, for example, that described in Gyuris et al., Cell 75:791-803 (1993)) to identify nucleic acids encoding the other polypeptide with which binding occurs, or to identify inhibitors of the binding interaction. Polypeptides involved in these binding interactions can also be used to screen for peptide or small molecule inhibitors or agonists of the binding interaction.

Competitive Binding Assays. Another type of suitable binding assay is a competitive binding assay. To illustrate, biological activity of a variant can be determined by assaying for the variant's ability to compete with the native polypeptide for binding to the candidate binding partner. Competitive binding assays can be performed by conventional methodology. Reagents that can be employed in competitive binding assays include radiolabeled HPR1 or HPR2 and intact cells expressing HPR1 and/or HPR2 (endogenous or recombinant) on the cell surface. For example, a radiolabeled soluble HPR1 or HPR2 fragment can be used to compete with a soluble HPR1 variant and/or a soluble HPR2 variant for binding to cell surface receptors. Instead of intact cells, one could

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substitute a soluble binding partner/Fc fusion polypeptide bound to a solid phase through the interaction of Polypeptide A or Polypeptide G (on the solid phase) with the Fc moiety. Chromatography columns that contain Polypeptide A and Polypeptide G include those available from Pharmacia Biotech, Inc., Piscataway, NJ.

Assays to Identify Modulators of Intracellular Signaling, Cell Proliferation, or Immune Cell Activity. The influence of HPR1 or HPR2 on intracellular signaling, cell proliferation, or immune cell activity can be manipulated to control these activities in target cells. For example, the disclosed HPR1 and HPR2 polypeptides, nucleic acids encoding the disclosed HPR1 and HPR2 polypeptides, or agonists or antagonists of such polypeptides can be administered to a cell or group of cells to induce, enhance, suppress, or arrest intracellular signaling or cell proliferation by the target cells. Identification of HPR1 and HPR2 polypeptides, agonists or antagonists that can be used in this manner can be carried out via a variety of assays known to those skilled in the art. Included in such assays are those that evaluate the ability of an HPR1 or HPR2 polypeptide to influence intracellular signaling, cell proliferation, or immune cell activity. Such an assay would involve, for example, the analysis of immune cell interaction in the presence of an HPR1 polypeptide and/or an HPR1 polypeptide. In such an assay, one would determine a rate of intracellular signaling or cell proliferation in the presence of the HPR1 and/or HPR2 polypeptide and then determine if such intracellular signaling or cell proliferation is altered in the presence of a candidate agonist or antagonist or another HPR1 or HPR2 polypeptide. Exemplary assays for this aspect of the invention include cytokine secretion assays, cell proliferation assays, and mixed lymphocyte reactions involving antigen presenting cells and T cells. These assays are well known to those skilled in the art.

In another aspect, the present invention provides a method of detecting the ability of a test compound to affect the intracellular signaling or cell proliferation activity of a cell. In this aspect, the method comprises: (1) contacting a first group of target cells with a test compound including an HPR1 polypeptide and/or an HPR2 polypeptide, or a fragment or fragments thereof, under conditions appropriate to the particular assay being used; (2) measuring the net rate of intracellular signaling or cell proliferation among the target cells; and (3) observing the net rate of intracellular signaling or cell proliferation among control cells contacting the HPR1 and/or HPR2 polypeptides or fragments thereof, in the absence of a test compound, under otherwise identical conditions as the first group of cells. In this embodiment, the net rate of intracellular signaling or cell proliferation in the control cells is compared to that of the cells treated with both a test compound and the HPR1 and/or HPR2 polypeptide(s). The comparison will provide a difference in the net rate of intracellular signaling or cell proliferation can be identified. The test compound can function as an effector by either activating or up-regulating, or by inhibiting or down-regulating, intracellular signaling or cell proliferation, and can be detected through this method.

<u>Cell Proliferation, Cell Death, Cell Differentiation, and Cell Adhesion Assays</u>. A polypeptide of the present invention may exhibit cytokine, cell proliferation (either inducing or inhibiting), or cell differentiation (either inducing or inhibiting) activity, or may induce production of other cytokines in

5 certain cell populations. Many polypeptide factors discovered to date have exhibited such activity in one or more factor-dependent cell proliferation assays, and hence the assays serve as a convenient confirmation of cell stimulatory activity. The activity of a polypeptide of the present invention is evidenced by any one of a number of routine factor-dependent cell proliferation assays for cell lines including, without limitation, 32D, DA2, DA1G, T10, B9, B9/11, BaF3, MC9/G, M+ (preB M+), 2E8, RB5, DA1, 123, T1165, HT2, CTLL2, TF-1, Mo7e and CMK. The activity of an HPR1 or HPR2 polypeptide of the invention may, among other means, be measured by the following methods:

Assays for T-cell or thymocyte proliferation include without limitation those described in: Current Protocols in Immunology, Coligan et al. eds, Greene Publishing Associates and Wiley-Interscience (pp. 3.1-3.19: In vitro assays for mouse lymphocyte function; Chapter 7: Immunologic studies in humans); Takai et al., J. Immunol. 137: 3494-3500, 1986; Bertagnolli et al., J. Immunol. 145: 1706-1712, 1990; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Bertagnolli, et al., J. Immunol. 149:3778-3783, 1992; Bowman et al., J. Immunol. 152: 1756-1761, 1994.

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Assays for cytokine production and/or proliferation of spleen cells, lymph node cells or thymocytes include, without limitation, those described in: Kruisbeek and Shevach, 1994, Polyclonal T cell stimulation, in *Current Protocols in Immunology*, Coligan *et al.* eds. Vol 1 pp. 3.12.1-3.12.14, John Wiley and Sons, Toronto; and Schreiber, 1994, Measurement of mouse and human interferon gamma in *Current Protocols in Immunology*, Coligan *et al.* eds. Vol 1 pp. 6.8.1-6.8.8, John Wiley and Sons, Toronto.

Assays for proliferation and differentiation of hematopoietic and lymphopoietic cells include, without limitation, those described in: Bottomly et al., 1991, Measurement of human and murine interleukin 2 and interleukin 4, in Current Protocols in Immunology, Coligan et al. eds. Vol 1 pp. 6.3.1-6.3.12, John Wiley and Sons, Toronto; deVries et al., J Exp Med 173: 1205-1211, 1991; Moreau et al., Nature 336:690-692, 1988; Greenberger et al., Proc Natl Acad Sci.USA 80: 2931-2938, 1983; Nordan, 1991, Measurement of mouse and human interleukin 6, in Current Protocols in Immunology Coligan et al. eds. Vol 1 pp. 6.6.1-6.6.5, John Wiley and Sons, Toronto; Smith et al., Proc Natl Acad Sci USA 83: 1857-1861, 1986; Bennett et al., 1991, Measurement of human interleukin 11, in Current Protocols in Immunology Coligan et al. eds. Vol 1 pp. 6.15.1 John Wiley and Sons, Toronto; Ciarletta et al., 1991, Measurement of mouse and human Interleukin 9, in Current Protocols in Immunology Coligan et al. eds. Vol 1 pp. 6.13.1, John Wiley and Sons, Toronto.

Assays for T-cell clone responses to antigens (which will identify, among others, polypeptides that affect APC-T cell interactions as well as direct T-cell effects by measuring proliferation and cytokine production) include, without limitation, those described in: Current Protocols in Immunology, Coligan et al. eds, Greene Publishing Associates and Wiley-Interscience (Chapter 3: In vitro assays for mouse lymphocyte function; Chapter 6: Cytokines and their cellular receptors; Chapter 7: Immunologic studies in humans); Weinberger et al., Proc Natl Acad Sci USA 77: 6091-6095, 1980; Weinberger et al., Eur. J. Immun. 11:405-411, 1981; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988

Assays for thymocyte or splenocyte cytotoxicity include, without limitation, those described in: Current Protocols in Immunology, Coligan et al. eds, Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Herrmann et al., Proc. Natl. Acad. Sci. USA 78:2488-2492, 1981; Herrmann et al., J. Immunol. 128:1968-1974, 1982; Handa et al., J. Immunol. 135:1564-1572, 1985; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 128:1968-1974, 1982; Handa et al., J. Immunol. 135:1564-1572, 1985; Takai et al., J. Immunol. 137:3494-3500, 1986; Bowmanet al., J. Virology 61:1992-1998; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Brown et al., J. Immunol. 153:3079-3092, 1994.

Assays for T-cell-dependent immunoglobulin responses and isotype switching (which will identify, among others, polypeptides that modulate T-cell dependent antibody responses and that affect Th1/Th2 profiles) include, without limitation, those described in: Maliszewski, J Immunol 144: 3028-3033, 1990; and Mond and Brunswick, 1994, Assays for B cell function: *in vitro* antibody production, in *Current Protocols in Immunology* Coligan *et al.* eds. Vol 1 pp. 3.8.1-3.8.16, John Wiley and Sons, Toronto.

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Mixed lymphocyte reaction (MLR) assays (which will identify, among others, polypeptides that generate predominantly Th1 and CTL responses) include, without limitation, those described in: Current Protocols in Immunology, Coligan et al. eds, Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., J. Immunol. 149:3778-3783, 1992.

Dendritic cell-dependent assays (which will identify, among others, polypeptides expressed by dendritic cells that activate naive T-cells) include, without limitation, those described in: Guery et al., J. Immunol 134:536-544, 1995; Inaba et al., J Exp Med 173:549-559, 1991; Macatonia et al., J Immunol 154:5071-5079, 1995; Porgador et al., J Exp Med 182:255-260, 1995; Nair et al., J Virology 67:4062-4069, 1993; Huang et al., Science 264:961-965, 1994; Macatonia et al., J Exp Med 169:1255-1264, 1989; Bhardwaj et al., J Clin Invest 94:797-807, 1994; and Inaba et al., J Exp Med 172:631-640,1990.

Assays for lymphocyte survival/apoptosis (which will identify, among others, polypeptides that prevent apoptosis after superantigen induction and polypeptides that regulate lymphocyte homeostasis) include, without limitation, those described in: Darzynkiewicz et al., Cytometry 13:795-808, 1992; Gorczyca et al., Leukemia 7:659-670, 1993; Gorczyca et al., Cancer Research 53:1945-1951, 1993; Itoh et al., Cell 66:233-243, 1991; Zacharchuk, J Immunol 145:4037-4045, 1990; Zamai et al., Cytometry 14:891-897, 1993; Gorczyca et al., International Journal of Oncology 1:639-648, 1992.

Assays for polypeptides that influence early steps of T-cell commitment and development include, without limitation, those described in: Antica et al., Blood 84:111-117, 1994; Fine et al., Cell Immunol 155:111-122, 1994; Galy et al., Blood 85:2770-2778, 1995; Toki et al., Proc Natl Acad Sci. USA 88:7548-7551, 1991

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Assays for embryonic stem cell differentiation (which will identify, among others, polypeptides that influence embryonic differentiation hematopoiesis) include, without limitation, those described in: Johansson et al. Cellular Biology 15:141-151, 1995; Keller et al., Molecular and Cellular Biology 13:473-486, 1993; McClanahan et al., Blood 81:2903-2915, 1993.

Assays for stem cell survival and differentiation (which will identify, among others, polypeptides that regulate lympho-hematopoiesis) include, without limitation, those described in: Methylcellulose colony forming assays, Freshney, 1994, In Culture of Hematopoietic Cells, Freshney et al. eds. pp. 265-268, Wiley-Liss, Inc., New York, NY; Hirayama et al., Proc. Natl. Acad. Sci. USA 89:5907-5911, 1992; Primitive hematopoietic colony forming cells with high proliferative potential, McNiece and Briddell, 1994, In Culture of Hematopoietic Cells, Freshney et al. eds. pp. 23-39, Wiley-Liss, Inc., New York, NY; Neben et al., Experimental Hematology 22:353-359, 1994; Ploemacher, 1994, Cobblestone area forming cell assay, In Culture of Hematopoietic Cells, Freshney et al. eds. pp. 1-21, Wiley-Liss, Inc., New York, NY; Spooncer et al., 1994, Long term bone marrow cultures in the presence of stromal cells, In Culture of Hematopoietic Cells, Freshney et al. eds. pp. 163-179, Wiley-Liss, Inc., New York, NY; Sutherland, 1994, Long term culture initiating cell assay, In Culture of Hematopoietic Cells, Freshney et al. eds. Vol pp. 139-162, Wiley-Liss, Inc., New York, NY.

Assay for hemostatic and thrombolytic activity include, without limitation, those described in: Linet et al., J. Clin. Pharmacol. 26:131-140, 1986; Burdick et al., Thrombosis Res. 45:413-419,1987; Humphrey et al., Fibrinolysis 5:71-79 (1991); Schaub, Prostaglandins 35:467-474, 1988.

Assays for receptor-ligand activity include without limitation those described in: Current Protocols in Immunology Coligan et al. eds, Greene Publishing Associates and Wiley-Interscience (Chapter 7.28, Measurement of cellular adhesion under static conditions 7.28.1-7.28.22), Takai et al., Proc. Natl. Acad. Sci. USA 84:6864-6868, 1987; Bierer et al., J. Exp. Med. 168:1145-1156, 1988; Rosenstein et al., J. Exp. Med. 169:149-160 1989; Stoltenborg et al., J. Immunol. Methods 175:59-68, 1994; Stitt et al., Cell 80:661-670, 1995.

## 30 Diagnostic and Other Uses of HPR1 and HPR2 Polypeptides and Nucleic Acids

The nucleic acids encoding the HPR1 and HPR2 polypeptides provided by the present invention can be used for numerous diagnostic or other useful purposes. The nucleic acids of the invention can be used to express recombinant polypeptide for analysis, characterization or therapeutic use; as markers for tissues in which the corresponding polypeptide is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in disease states); as molecular weight markers on Southern gels; as chromosome markers or tags (when labeled) to identify chromosomes or to map related gene positions; to compare with endogenous DNA sequences in patients to identify potential genetic disorders; as probes to hybridize and thus discover novel, related DNA sequences; as a source of information to derive PCR primers for genetic fingerprinting; as a probe to "subtract-out" known sequences in the process of discovering other novel nucleic acids; for selecting and making oligomers for attachment to a "gene chip" or other support, including for examination of expression patterns; to raise anti-polypeptide antibodies using DNA immunization

techniques; as an antigen to raise anti-DNA antibodies or elicit another immune response, and. for gene therapy. Uses of HPR1 and HPR2 polypeptides and fragmented polypeptides include, but are not limited to, the following: purifying polypeptides and measuring the activity thereof; delivery agents; therapeutic and research reagents; molecular weight and isoelectric focusing markers; controls for peptide fragmentation; identification of unknown polypeptides; and preparation of antibodies. Any or all nucleic acids suitable for these uses are capable of being developed into reagent grade or kit format for commercialization as products. Methods for performing the uses listed above are well known to those skilled in the art. References disclosing such methods include without limitation "Molecular Cloning: A Laboratory Manual", 2d ed., Cold Spring Harbor Laboratory Press, Sambrook, J., E. F. Fritsch and T. Maniatis eds., 1989, and "Methods in Enzymology: Guide to Molecular Cloning Techniques", Academic Press, Berger, S. L. and A. R. Kimmel eds., 1987

Probes and Primers. Among the uses of the disclosed HPR1 and HPR2 nucleic acids, and combinations of fragments thereof, is the use of fragments as probes or primers. Such fragments generally comprise at least about 17 contiguous nucleotides of a DNA sequence. In other embodiments, a DNA fragment comprises at least 30, or at least 60, contiguous nucleotides of a DNA sequence. The basic parameters affecting the choice of hybridization conditions and guidance for devising suitable conditions are set forth by Sambrook et al., 1989 and are described in detail above. Using knowledge of the genetic code in combination with the amino acid sequences set forth above, sets of degenerate oligonucleotides can be prepared. Such oligonucleotides are useful as primers, e.g., in polymerase chain reactions (PCR), whereby DNA fragments are isolated and amplified. In certain embodiments, degenerate primers can be used as probes for non-human genetic libraries. Such libraries would include but are not limited to cDNA libraries, genomic libraries, and even electronic EST (express sequence tag) or DNA libraries. Homologous sequences identified by this method would then be used as probes to identify non-human HPR1 and HPR2 homologues.

Chromosome Mapping. The nucleic acids encoding HPR1 and HPR2 polypeptides, and the disclosed fragments and combinations of these nucleic acids, can be used by those skilled in the art using well-known techniques to identify the human chromosome to which these nucleic acids map. Useful techniques include, but are not limited to, using the sequence or portions, including oligonucleotides, as a probe in various well-known techniques such as radiation hybrid mapping (high resolution), in situ hybridization to chromosome spreads (moderate resolution), and Southern blot hybridization to hybrid cell lines containing individual human chromosomes (low resolution). Alternatively, the genomic sequences corresponding to nucleic acids encoding a cytokine polypeptide of the invention are mapped by comparison to sequences in public and proprietary databases, such as GenBank (ncbi.nlm.nih.gov/BLAST), Locuslink (ncbi.nlm.nih.gov:80/LocusLink/), Unigene (ncbi.nlm.nih.gov/cgi-bin/UniGene), AceView (ncbi.nlm.nih.gov/AceView), Gene Map Viewer (ncbi.nlm.nih.gov/genemap), Online Mendelian Inheritance in Man (OMIM) (ncbi.nlm.nih.gov/Omim), and proprietary databases such as the Celera Discovery System (celera.com). These computer analyses of available genomic sequence information can provide the identification of the specific chromosomal

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location of human and/or murine genomic sequences corresponding to sequences encoding HPR1 or HPR2 polypeptides of the invention, and the unique genetic mapping relationships between HPR1 or HPR2 genomic sequences and the genetic map locations of known human genetic disorders

Diagnostics and Gene Therapy. The nucleic acids encoding HPR1 and HPR2 polypeptides, and the disclosed fragments and combinations of these nucleic acids can be used by one skilled in the art using well-known techniques to analyze abnormalities associated with the genes corresponding to these polypeptides. This enables one to distinguish conditions in which this marker is rearranged or deleted. In addition, nucleic acids of the invention or a fragment thereof can be used as a positional marker to map other genes of unknown location. The DNA can be used in developing treatments for any disorder mediated (directly or indirectly) by defective, or insufficient amounts of, the genes corresponding to the nucleic acids of the invention. Disclosure herein of native nucleotide sequences permits the detection of defective genes, and the replacement thereof with normal genes. Defective genes can be detected in *in vitro* diagnostic assays, and by comparison of a native nucleotide sequence disclosed herein with that of a gene derived from a person suspected of harboring a defect in this gene.

Methods of Screening for Binding Partners. The HPR1 and HPR2 polypeptides of the invention each can be used as reagents in methods to screen for or identify binding partners. For example, the HPR1 and HPR2 polypeptides can be attached to a solid support material and may bind to their binding partners in a manner similar to affinity chromatography. In particular embodiments, a polypeptide is attached to a solid support by conventional procedures. As one example, chromatography columns containing functional groups that will react with functional groups on amino acid side chains of polypeptides are available (Pharmacia Biotech, Inc., Piscataway, NJ). In an alternative, a polypeptide/Fc polypeptide (as discussed above) is attached to Protein A- or Protein Gcontaining chromatography columns through interaction with the Fc moiety. The HPR1 and HPR2 polypeptides also find use in identifying cells that express a binding partner on the cell surface. Polypeptides are bound to a solid phase such as a column chromatography matrix or a similar suitable substrate. For example, magnetic microspheres can be coated with the polypeptides and held in an incubation vessel through a magnetic field. Suspensions of cell mixtures containing potential bindingpartner-expressing cells are contacted with the solid phase having the polypeptides thereon. Cells expressing the binding partner on the cell surface bind to the fixed polypeptides, and unbound cells are washed away. Alternatively, HPR1 and HPR2 polypeptides can be conjugated to a detectable moiety, then incubated with cells to be tested for binding partner expression. After incubation, unbound labeled matter is removed and the presence or absence of the detectable moiety on the cells is determined. In a further alternative, mixtures of cells suspected of expressing the binding partner are incubated with biotinylated polypeptides. Incubation periods are typically at least one hour in duration to ensure sufficient binding. The resulting mixture then is passed through a column packed with avidin-coated beads, whereby the high affinity of biotin for avidin provides binding of the desired cells to the beads. Procedures for using avidin-coated beads are known (see Berenson, et al. J. Cell. Biochem., 10D:239, 1986). Washing to remove unbound material, and the release of the bound cells,

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are performed using conventional methods. In some instances, the above methods for screening for or identifying binding partners may also be used or modified to isolate or purify such binding partner molecules or cells expressing them.

Measuring Biological Activity. HPR1 and HPR2 polypeptides also find use in measuring the biological activity of HPR1-binding and/or HPR2-binding polypeptides in terms of their binding affinity. The polypeptides thus can be employed by those conducting "quality assurance" studies, e.g., to monitor shelf life and stability of polypeptide under different conditions. For example, the polypeptides can be employed in a binding affinity study to measure the biological activity of a binding partner polypeptide that has been stored at different temperatures, or produced in different cell types. The polypeptides also can be used to determine whether biological activity is retained after modification of a binding partner polypeptide (e.g., chemical modification, truncation, mutation, etc.). The binding affinity of the modified polypeptide is compared to that of an unmodified binding polypeptide to detect any adverse impact of the modifications on biological activity of the binding polypeptide. The biological activity of a binding polypeptide thus can be ascertained before it is used in a research study, for example.

Carriers and Delivery Agents. The polypeptides also find use as carriers for delivering agents attached thereto to cells bearing identified binding partners. The polypeptides thus can be used to deliver diagnostic or therapeutic agents to such cells (or to other cell types found to express binding partners on the cell surface) in in vitro or in vivo procedures. Detectable (diagnostic) and therapeutic agents that can be attached to a polypeptide include, but are not limited to, toxins, other cytotoxic agents, drugs, radionuclides, chromophores, enzymes that catalyze a colorimetric or fluorometric reaction, and the like, with the particular agent being chosen according to the intended application. Among the toxins are ricin, abrin, diphtheria toxin, Pseudomonas aeruginosa exotoxin A, ribosomal inactivating polypeptides, mycotoxins such as trichothecenes, and derivatives and fragments (e.g., single chains) thereof. Radionuclides suitable for diagnostic use include, but are not limited to, <sup>123</sup>I, <sup>131</sup>I, <sup>99m</sup>Tc, <sup>111</sup>In, and <sup>76</sup>Br. Examples of radionuclides suitable for therapeutic use are <sup>131</sup>I, <sup>211</sup>At, <sup>77</sup>Br, <sup>186</sup>Re, <sup>188</sup>Re, <sup>212</sup>Pb, <sup>212</sup>Bi, <sup>109</sup>Pd, <sup>64</sup>Cu, and <sup>67</sup>Cu. Such agents can be attached to the polypeptide by any suitable conventional procedure. The polypeptide comprises functional groups on amino acid side chains that can be reacted with functional groups on a desired agent to form covalent bonds, for example. Alternatively, the polypeptide or agent can be derivatized to generate or attach a desired reactive functional group. The derivatization can involve attachment of one of the bifunctional coupling reagents available for attaching various molecules to polypeptides (Pierce Chemical Company, Rockford, Illinois). A number of techniques for radiolabeling polypeptides are known. Radionuclide metals can be attached to polypeptides by using a suitable bifunctional chelating agent, for example. Conjugates comprising polypeptides and a suitable diagnostic or therapeutic agent (preferably covalently linked) are thus prepared. The conjugates are administered or otherwise employed in an amount appropriate for the particular application.

#### 5 Treating Diseases Using HPR1 and/or HPR2 Polypeptides and Antagonists Thereof

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It is anticipated that the HPR1 and HPR2 polypeptides, fragments, variants, antagonists, agonists, antibodies, and binding partners of the invention will be useful for treating medical conditions and diseases including, but not limited to, cell proliferation, metabolic, and reproductive hormone related conditions as described further herein. The therapeutic molecule or molecules to be used will depend on the etiology of the condition to be treated and the biological pathways involved, and variants, fragments, and binding partners of HPR1 and/or HPR2 polypeptides may have effects similar to or different from HPR1 or HPR2 polypeptides. For example, an antagonist of the ligand-binding activity of HPR1 and/or HPR2 polypeptides may be selected for treatment of conditions involving ligand-binding activity, but a particular fragment of a given HPR1 or HPR2 polypeptide may also act as an effective dominant negative antagonist of that activity. Therefore, in the following paragraphs "HPR1 and HPR2 polypeptides or antagonists" refers to all HPR1 and HPR2 polypeptides, fragments, variants, antagonists, agonists, antibodies, and binding partners etc. of the invention, and it is understood that a specific molecule or molecules can be selected from those provided as embodiments of the invention by individuals of skill in the art, according to the biological and therapeutic considerations described herein.

Provided herein are methods for using HPR1 and HPR2 polypeptides or antagonists, compositions or combination therapies to treat various hematologic and oncologic disorders. For example, HPR1 and HPR2 polypeptides or antagonists are used to treat various forms of cancer, including acute myelogenous leukemia, Epstein-Barr virus-positive nasopharyngeal carcinoma, glioma, colon, stomach, prostate, renal cell, cervical and ovarian cancers, lung cancer (SCLC and NSCLC), including cancer-associated cachexia, fatigue, asthenia, paraneoplastic syndrome of cachexia and hypercalcemia. Additional diseases treatable with the subject HPR1 and HPR2 polypeptides or antagonists, compositions or combination therapies are solid tumors, including sarcoma, osteosarcoma, and carcinoma, such as adenocarcinoma (for example, breast cancer) and squamous cell carcinoma. In addition, the subject compounds, compositions or combination therapies are useful for treating leukemia, including acute myelogenous leukemia, chronic or acute lymphoblastic leukemia and hairy cell leukemia. Other malignancies with invasive metastatic potential can be treated with the subject compounds, compositions and combination therapies, including multiple myeloma. In addition, the disclosed HPR1 and HPR2 polypeptides or antagonists, compositions and combination therapies can be used to treat anemias and hematologic disorders, including anemia of chronic disease, aplastic anemia, including Fanconi's aplastic anemia; idiopathic thrombocytopenic purpura (ITP); myelodysplastic syndromes (including refractory anemia, refractory anemia with ringed sideroblasts, refractory anemia with excess blasts, refractory anemia with excess blasts in transformation); myelofibrosis/myeloid metaplasia; and sickle cell vasocclusive crisis.

Various lymphoproliferative disorders also are treatable with the disclosed HPR1 and HPR2 polypeptides or antagonists, compositions or combination therapies. These include, but are not limited to autoimmune lymphoproliferative syndrome (ALPS), chronic lymphoblastic leukemia, hairy cell

5 Ieukemia, chronic lymphatic leukemia, peripheral T-cell lymphoma, small lymphocytic lymphoma, mantle cell lymphoma, follicular lymphoma, Burkitt's lymphoma, Epstein-Barr virus-positive T cell lymphoma, histiocytic lymphoma, Hodgkin's disease, diffuse aggressive lymphoma, acute lymphatic leukemias, T gamma lymphoproliferative disease, cutaneous B cell lymphoma, cutaneous T cell lymphoma (i.e., mycosis fungoides) and Sézary syndrome.

In addition, the subject invention provides HPR1 and HPR2 polypeptides or antagonists, compositions and combination therapies for the treatment of non-arthritic medical conditions of the bones and joints. This encompasses osteoclast disorders that lead to bone loss, such as but not limited to osteoporosis, including post-menopausal osteoporosis, periodontitis resulting in tooth loosening or loss, and prosthesis loosening after joint replacement (generally associated with an inflammatory response to wear debris). This latter condition also is called "orthopedic implant osteolysis." Another condition treatable by administering HPR1 and HPR2 polypeptides or antagonists, is temporal mandibular joint dysfunction (TMI).

The disclosed HPR1 and HPR2 polypeptides or antagonists, compositions and combination therapies furthermore are useful for treating neurodegenerative conditions such as acute polyneuropathy; anorexia nervosa; Bell's palsy; chronic fatigue syndrome; transmissible dementia, including Creutzfeld-Jacob disease; demyelinating neuropathy; Guillain-Barre syndrome; vertebral disc disease; Gulf war syndrome; myasthenia gravis; chronic neuronal degeneration; and stroke, including cerebral ischemic diseases.

# 25 Administration of HPR1 and HPR2 Polypeptides and Antagonists Thereof

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This invention provides compounds, compositions, and methods for treating a patient, preferably a mammalian patient, and most preferably a human patient, who is suffering from a medical disorder, and in particular an HPR1- or HPR2-mediated disorder. Such HPR1- or HPR2-mediated disorders include conditions caused (directly or indirectly) or exacerbated by binding between HPR1 and/or HPR2 and a binding partner. For purposes of this disclosure, the terms "illness," "disease," "medical condition," "abnormal condition" and the like are used interchangeably with the term "medical disorder." The terms "treat", "treating", and "treatment" used herein includes curative, preventative (e.g., prophylactic) and palliative or ameliorative treatment. For such therapeutic uses, HPR1 and HPR2 polypeptides and fragments, HPR1 and HPR2 nucleic acids encoding the HPR1 and HPR2 polypeptides, and/or agonists or antagonists of the HPR1 and/or HPR2 polypeptides such as antibodies can be administered to the patient in need through well-known means. Compositions of the present invention can contain a polypeptide in any form described herein, such as native polypeptides, variants, derivatives, oligomers, and biologically active fragments. In particular embodiments, the composition comprises a soluble polypeptide or an oligomer comprising soluble HPR1 and/or HPR2 polypeptides.

<u>Therapeutically Effective Amount</u>. In practicing the method of treatment or use of the present invention, a therapeutically effective amount of a therapeutic agent of the present invention is

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administered to a patient having a condition to be treated, preferably to treat or ameliorate diseases associated with the activity of an HPR1 and/or HPR2 polypeptide. "Therapeutic agent" includes without limitation any of the HPR1 or HPR2 polypeptides, fragments, and variants; nucleic acids encoding the HPR1 and HPR2 polypeptides, fragments, and variants; agonists or antagonists of the HPR1 and HPR2 polypeptides such as antibodies; HPR1 and/or HPR2 polypeptide binding partners; complexes formed from the HPR1 and/or HPR2 polypeptides, fragments, variants, and binding partners, etc. As used herein, the term "therapeutically effective amount" means the total amount of each therapeutic agent or other active component of the pharmaceutical composition or method that is sufficient to show a meaningful patient benefit, i.e., treatment, healing, prevention or amelioration of the relevant medical condition, or an increase in rate of treatment, healing, prevention or amelioration of such conditions. When applied to an individual therapeutic agent or active ingredient, administered alone, the term refers to that ingredient alone. When applied to a combination, the term refers to combined amounts of the ingredients that result in the therapeutic effect, whether administered in combination, serially or simultaneously. As used herein, the phrase "administering a therapeutically effective amount" of a therapeutic agent means that the patient is treated with said therapeutic agent in an amount and for a time sufficient to induce an improvement, and preferably a sustained improvement, in at least one indicator that reflects the severity of the disorder. An improvement is considered "sustained" if the patient exhibits the improvement on at least two occasions separated by one or more weeks. The degree of improvement is determined based on signs or symptoms, and determinations can also employ questionnaires that are administered to the patient, such as quality-oflife questionnaires. Various indicators that reflect the extent of the patient's illness can be assessed for determining whether the amount and time of the treatment is sufficient. The baseline value for the chosen indicator or indicators is established by examination of the patient prior to administration of the first dose of the therapeutic agent. Preferably, the baseline examination is done within about 60 days of administering the first dose. If the therapeutic agent is being administered to treat acute symptoms, the first dose is administered as soon as practically possible after the injury has occurred. Improvement is induced by administering therapeutic agents such as HPR1 and/or HPR2 polypeptides or antagonists until the patient manifests an improvement over baseline for the chosen indicator or indicators. In treating chronic conditions, this degree of improvement is obtained by repeatedly administering this medicament over a period of at least a month or more, e.g., for one, two, or three months or longer, or indefinitely. A period of one to six weeks, or even a single dose, often is sufficient for treating injuries or acute conditions. Although the extent of the patient's illness after treatment may appear improved according to one or more indicators, treatment may be continued indefinitely at the same level or at a reduced dose or frequency. Once treatment has been reduced or discontinued, it later may be resumed at the original level if symptoms should reappear.

<u>Dosing.</u> One skilled in the pertinent art will recognize that suitable dosages will vary, depending upon such factors as the nature and severity of the disorder to be treated, the patient's body weight, age, general condition, and prior illnesses and/or treatments, and the route of administration.

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Preliminary doses can be determined according to animal tests, and the scaling of dosages for human administration is performed according to art-accepted practices such as standard dosing trials. For example, the therapeutically effective dose can be estimated initially from cell culture assays. The dosage will depend on the specific activity of the compound and can be readily determined by routine experimentation. A dose can be formulated in animal models to achieve a circulating plasma concentration range that includes the IC50 (i.e., the concentration of the test compound which achieves a half-maximal inhibition of symptoms) as determined in cell culture, while minimizing toxicities. Such information can be used to more accurately determine useful doses in humans. Ultimately, the attending physician will decide the amount of polypeptide of the present invention with which to treat each individual patient. Initially, the attending physician will administer low doses of polypeptide of the present invention and observe the patient's response. Larger doses of polypeptide of the present invention can be administered until the optimal therapeutic effect is obtained for the patient, and at that point the dosage is not increased further. It is contemplated that the various pharmaceutical compositions used to practice the method of the present invention should contain about 0.01 ng to about 100 mg (preferably about 0.1 ng to about 10 mg, more preferably about 0.1 microgram to about 1 mg) of polypeptide of the present invention per kg body weight. In one embodiment of the invention, HPR1 and/or HPR2 polypeptides or antagonists are administered one time per week to treat the various medical disorders disclosed herein, in another embodiment is administered at least two times per week, and in another embodiment is administered at least three times per week. If injected, the effective amount of HPR1 or HPR2 polypeptides or antagonists per adult dose ranges from 1-20 mg/m<sup>2</sup>, and preferably is about 5-12 mg/m<sup>2</sup>. Alternatively, a flat dose can be administered, whose amount may range from 5-100 mg/dose. Exemplary dose ranges for a flat dose to be administered by subcutaneous injection are 5-25 mg/dose, 25-50 mg/dose and 50-100 mg/dose. In one embodiment of the invention, the various indications described below are treated by administering a preparation acceptable for injection containing HPR1 and/or HPR2 polypeptides or antagonists at 25 mg/dose, or alternatively, containing 50 mg per dose. The 25 mg or 50 mg dose can be administered repeatedly, particularly for chronic conditions. If a route of administration other than injection is used, the dose is appropriately adjusted in accord with standard medical practices. In many instances, an improvement in a patient's condition will be obtained by injecting a dose of about 25 mg of HPR1 or HPR2 polypeptides or antagonists one to three times per week over a period of at least three weeks, or a dose of 50 mg of HPR1 or HPR2 polypeptides or antagonists one or two times per week for at least three weeks, though treatment for longer periods may be necessary to induce the desired degree of improvement. For incurable chronic conditions, the regimen can be continued indefinitely, with adjustments being made to dose and frequency if such are deemed necessary by the patient's physician. The foregoing doses are examples for an adult patient who is a person who is 18 years of age or older. For pediatric patients (age 4-17), a suitable regimen involves the subcutaneous injection of 0.4 mg/kg, up to a maximum dose of 25 mg of HPR1 or HPR2 polypeptides or antagonists, administered by subcutaneous injection one or more times per week. If an antibody against an HPR1 and/or HPR2 polypeptide is used as the HPR1

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and/or HPR2 polypeptide antagonist, a preferred dose range is 0.1 to 20 mg/kg, and more preferably is 1-10 mg/kg. Another preferred dose range for an anti-HPR1 polypeptide and/or anti-HPR2 polypeptide antibody is 0.75 to 7.5 mg/kg of body weight. Humanized antibodies are preferred, that is, antibodies in which only the antigen-binding portion of the antibody molecule is derived from a non-human source. Such antibodies can be injected or administered intravenously.

Formulations. Compositions comprising an effective amount of an HPR1 and/or HPR2 polypeptide of the present invention (from whatever source derived, including without limitation from recombinant and non-recombinant sources), in combination with other components such as a physiologically acceptable diluent, carrier, or excipient, are provided herein. The term "pharmaceutically acceptable" means a non-toxic material that does not interfere with the effectiveness of the biological activity of the active ingredient(s). Formulations suitable for administration include aqueous and non-aqueous sterile injection solutions which can contain anti-oxidants, buffers, bacteriostats and solutes which render the formulation isotonic with the blood of the recipient; and aqueous and non-aqueous sterile suspensions which can include suspending agents or thickening The polypeptides can be formulated according to known methods used to prepare agents. pharmaceutically useful compositions. They can be combined in admixture, either as the sole active material or with other known active materials suitable for a given indication, with pharmaceutically acceptable diluents, preservatives, emulsifiers, solubilizers, adjuvants and/or carriers. Suitable formulations for pharmaceutical compositions include those described in Remington's Pharmaceutical Sciences, 16th ed. 1980, Mack Publishing Company, Easton, PA. In addition, such compositions can be complexed with polyethylene glycol (PEG), metal ions, or incorporated into polymeric compounds such as polyacetic acid, polyglycolic acid, hydrogels, dextran, etc., or incorporated into liposomes, microemulsions, micelles, unilamellar or multilamellar vesicles, erythrocyte ghosts or spheroblasts. Suitable lipids for liposomal formulation include, without limitation, monoglycerides, diglycerides, sulfatides, lysolecithin, phospholipids, saponin, bile acids, and the like. Preparation of such liposomal formulations is within the level of skill in the art, as disclosed, for example, in U.S. Pat. No. 4,235,871; U.S. Pat. No. 4,501,728; U.S. Pat. No. 4,837,028; and U.S. Pat. No. 4,737,323. Such compositions will influence the physical state, solubility, stability, rate of in vivo release, and rate of in vivo clearance, and are thus chosen according to the intended application, so that the characteristics of the carrier will depend on the selected route of administration. In one preferred embodiment of the invention, sustained-release forms of HPR1 and/or HPR2 polypeptides are used. Sustained-release forms suitable for use in the disclosed methods include, but are not limited to, HPR1 and/or HPR2 polypeptides that are encapsulated in a slowly-dissolving biocompatible polymer (such as the alginate microparticles described in U.S. No. 6,036,978), admixed with such a polymer (including topically applied hydrogels), and or encased in a biocompatible semi-permeable implant.

<u>Combinations of Therapeutic Compounds.</u> An HPR1 or HPR2 polypeptide of the present invention may be active in multimers (e.g., heterodimers or homodimers) or complexes with itself or other polypeptides. As a result, pharmaceutical compositions of the invention may comprise a

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polypeptide of the invention in such multimeric or complexed form. The pharmaceutical composition of the invention may be in the form of a complex of the polypeptide(s) of present invention along with polypeptide or peptide antigens. The invention further includes the administration of HPR1 and/or HPR2 polypeptides or antagonists concurrently with one or more other drugs that are administered to the same patient in combination with the HPR1 and/or HPR2 polypeptides or antagonists, each drug being administered according to a regimen suitable for that medicament. "Concurrent administration" encompasses simultaneous or sequential treatment with the components of the combination, as well as regimens in which the drugs are alternated, or wherein one component is administered long-term and the other(s) are administered intermittently. Components can be administered in the same or in separate compositions, and by the same or different routes of administration. Examples of components that can be included in the pharmaceutical composition of the invention are: cytokines, lymphokines, or other hematopoietic factors such as M-CSF, GM-CSF, TNF, IL-1, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-11, IL-12, IL-13, IL-14, IL-15, IL-17, IL-18, IL-23, IFN, TNF0, TNF1, TNF2, G-CSF, Meg-CSF, thrombopoietin, stem cell factor, and erythropoietin. The pharmaceutical composition can further contain other agents which either enhance the activity of the polypeptide or compliment its activity or use in treatment. Such additional factors and/or agents may be included in the pharmaceutical composition to produce a synergistic effect with polypeptide of the invention, or to minimize side effects. Additional examples of drugs to be administered concurrently include but are not limited to antivirals, antibiotics, analgesics, corticosteroids, antagonists of inflammatory cytokines, non-steroidal anti-inflammatories, pentoxifylline, thalidomide, and disease-modifying antirheumatic drugs (DMARDs) such as azathioprine, cyclophosphamide, cyclosporine, hydroxychloroquine sulfate, methotrexate, leflunomide, minocycline, penicillamine, sulfasalazine and gold compounds such as oral gold, gold sodium thiomalate, and aurothioglucose. Additionally, HPR1 and/or HPR2 polypeptides or antagonists can be combined with a second HPR1 and/or HPR2 polypeptide/antagonist, including an antibody against an HPR1 and/or HPR2 polypeptide, or an HPR1 polypeptide-derived peptide or HPR2 polypeptide-derived peptide that acts as a competitive inhibitor of native HPR1 and/or HPR2 polypeptides.

Routes of Administration. Any efficacious route of administration may be used to therapeutically administer HPR1 and HPR2 polypeptides or antagonists thereof, including those compositions comprising nucleic acids. Parenteral administration includes injection, for example, via intra-articular, intravenous, intramuscular, intralesional, intraperitoneal or subcutaneous routes by bolus injection or by continuous infusion., and also includes localized administration, e.g., at a site of disease or injury. Other suitable means of administration include sustained release from implants; aerosol inhalation and/or insufflation.; eyedrops; vaginal or rectal suppositories; buccal preparations; oral preparations, including pills, syrups, lozenges or chewing gum; and topical preparations such as lotions, gels, sprays, ointments or other suitable techniques. Alternatively, polypeptideaceous HPR1 and HPR2 polypeptides or antagonists may be administered by implanting cultured cells that express the polypeptide, for example, by implanting cells that express HPR1 and/or HPR2 polypeptides or

antagonists. Cells may also be cultured ex vivo in the presence of polypeptides of the present invention in order to proliferate or to produce a desired effect on or activity in such cells. Treated cells can then be introduced in vivo for therapeutic purposes. In another embodiment, the patient's own cells are induced to produce HPR1 and/or HPR2 polypeptides or antagonists by transfection *in vivo* or *ex vivo* with a DNA that encodes HPR1 and/or HPR2 polypeptides or antagonists. This DNA can be introduced into the patient's cells, for example, by injecting naked DNA or liposome-encapsulated DNA that encodes HPR1 and/or HPR2 polypeptides or antagonists, or by other means of transfection. Nucleic acids of the invention can also be administered to patients by other known methods for introduction of nucleic acid into a cell or organism (including, without limitation, in the form of viral vectors or naked DNA). When HPR1 and/or HPR2 polypeptides or antagonists are administered in combination with one or more other biologically active compounds, these can be administered by the same or by different routes, and can be administered simultaneously, separately or sequentially.

Oral Administration. When a therapeutically effective amount of polypeptide of the present invention is administered orally, polypeptide of the present invention will be in the form of a tablet, capsule, powder, solution or elixir. When administered in tablet form, the pharmaceutical composition of the invention can additionally contain a solid carrier such as a gelatin or an adjuvant. The tablet, capsule, and powder contain from about 5 to 95% polypeptide of the present invention, and preferably from about 25 to 90% polypeptide of the present invention. When administered in liquid form, a liquid carrier such as water, petroleum, oils of animal or plant origin such as peanut oil, mineral oil, soybean oil, or sesame oil, or synthetic oils can be added. The liquid form of the pharmaceutical composition can further contain physiological saline solution, dextrose or other saccharide solution, or glycols such as ethylene glycol, propylene glycol or polyethylene glycol. When administered in liquid form, the pharmaceutical composition contains from about 0.5 to 90% by weight of polypeptide of the present invention, and preferably from about 1 to 50% polypeptide of the present invention.

Intravenous Administration. When a therapeutically effective amount of polypeptide of the present invention is administered by intravenous, cutaneous or subcutaneous injection, polypeptide of the present invention will be in the form of a pyrogen-free, parenterally acceptable aqueous solution. The preparation of such parenterally acceptable polypeptide solutions, having due regard to pH, isotonicity, stability, and the like, is within the skill in the art. A preferred pharmaceutical composition for intravenous, cutaneous, or subcutaneous injection should contain, in addition to polypeptide of the present invention, an isotonic vehicle such as Sodium Chloride Injection, Ringer's Injection, Dextrose Injection, Dextrose and Sodium Chloride Injection, Lactated Ringer's Injection, or other vehicle as known in the art. The pharmaceutical composition of the present invention can also contain stabilizers, preservatives, buffers, antioxidants, or other additives known to those of skill in the art. The duration of intravenous therapy using the pharmaceutical composition of the present invention will vary, depending on the severity of the disease being treated and the condition and potential idiosyncratic response of each individual patient. It is contemplated that the duration of each application of the polypeptide of the present invention will be in the range of 12 to 24 hours of continuous intravenous

administration. Ultimately the attending physician will decide on the appropriate duration of intravenous therapy using the pharmaceutical composition of the present invention.

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Bone and Tissue Administration. For compositions of the present invention which are useful for bone, cartilage, tendon or ligament disorders, the therapeutic method includes administering the composition topically, systematically, or locally as an implant or device. When administered, the therapeutic composition for use in this invention is, of course, in a pyrogen-free, physiologically acceptable form. Further, the composition can desirably be encapsulated or injected in a viscous form for delivery to the site of bone, cartilage or tissue damage. Topical administration may be suitable for wound healing and tissue repair. Therapeutically useful agents other than a polypeptide of the invention which can also optionally be included in the composition as described above, can alternatively or additionally, be administered simultaneously or sequentially with the composition in the methods of the invention. Preferably for bone and/or cartilage formation, the composition would include a matrix capable of delivering the polypeptide-containing composition to the site of bone and/or cartilage damage, providing a structure for the developing bone and cartilage and optimally capable of being resorbed into the body. Such matrices can be formed of materials presently in use for other implanted medical applications. The choice of matrix material is based on biocompatibility, biodegradability, mechanical properties, cosmetic appearance and interface properties. The particular application of the compositions will define the appropriate formulation. Potential matrices for the compositions can be biodegradable and chemically defined calcium sulfate, tricalciumphosphate, hydroxyapatite, polylactic acid, polyglycolic acid and polyanhydrides. Other potential materials are biodegradable and biologically well-defined, such as bone or dermal collagen. Further matrices are comprised of pure polypeptides or extracellular matrix components. Other potential matrices are nonbiodegradable and chemically defined, such as sintered hydroxapatite, bioglass, aluminates, or other ceramics Matrices can be comprised of combinations of any of the above mentioned types of material, such as polylactic acid and hydroxyapatite or collagen and tricalciumphosphate. Presently preferred is a 50:50 (mole weight) copolymer of lactic acid and glycolic acid in the form of porous particles having diameters ranging from 150 to 800 microns. In some applications, it will be useful to utilize a sequestering agent, such as carboxymethyl cellulose or autologous blood clot, to prevent the polypeptide compositions from disassociating from the matrix. A preferred family of sequestering agents is cellulosic materials such as alkylcelluloses, the most preferred being cationic salts of carboxymethylcellulose (CMC). Other preferred sequestering agents include hyaluronic acid, sodium alginate, poly(ethylene glycol), polyoxyethylene oxide, carboxyvinyl polymer and poly(vinyl alcohol). The amount of sequestering agent useful herein is 0.5-20 wt %, preferably 1-10 wt % based on total formulation weight, which represents the amount necessary to prevent desorbtion of the polypeptide from the polymer matrix and to provide appropriate handling of the composition, yet not so much that the progenitor cells are prevented from infiltrating the matrix, thereby providing the polypeptide the opportunity to assist the osteogenic activity of the progenitor cells. In further compositions, polypeptides of the invention can be combined with other agents beneficial to the treatment of the bone

and/or cartilage defect, wound, or tissue in question. Progress can be monitored by periodic assessment of tissue/bone growth and/or repair, for example, X-rays, histomorphometric determinations and tetracycline labeling.

Veterinary Uses. In addition to human patients, HPR1 and HPR2 polypeptides and antagonists are useful in the treatment of disease conditions in non-human animals, such as pets (dogs, cats, birds, primates, etc.), domestic farm animals (horses cattle, sheep, pigs, birds, etc.), or any animal that suffers from a TNFα-mediated inflammatory or arthritic condition. In such instances, an appropriate dose can be determined according to the animal's body weight. For example, a dose of 0.2-1 mg/kg may be used. Alternatively, the dose is determined according to the animal's surface area, an exemplary dose ranging from 0.1-20 mg/m², or more preferably, from 5-12 mg/m². For small animals, such as dogs or cats, a suitable dose is 0.4 mg/kg. In a preferred embodiment, HPR1 and/or HPR2 polypeptides or antagonists (preferably constructed from genes derived from the same species as the patient), are administered by injection or other suitable route one or more times per week until the animal's condition is improved, or they can be administered indefinitely.

Manufacture of Medicaments. The present invention also relates to the use of HPR1 and HPR2 polypeptides, fragments, and variants; nucleic acids encoding the HPR1 or HPR2 polypeptides, fragments, and variants; agonists or antagonists of the HPR1 and/or HPR2 polypeptides such as antibodies; HPR1 and/or HPR2 polypeptide binding partners; complexes formed from the HPR1 and/or HPR2 polypeptides, fragments, variants, and binding partners, etc, in the manufacture of a medicament for the prevention or therapeutic treatment of each medical disorder disclosed herein.

25 EXAMPLES

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The following examples are intended to illustrate particular embodiments and not to limit the scope of the invention.

#### EXAMPLE 1

#### A. Identification of HPR1, a New Member of the Human Hematopoietin Receptor Family

A data set was received from Celera Genomics (Rockville, Maryland) containing a listing of amino acid sequences predicted to be encoded by the human genome. This data set was searched with a BLAST algorithm to identify hematopoietin receptor family polypeptides. Several amino acid sequences, including two overlapping amino acid sequences (SEQ ID NO: 1 and SEQ ID NO:2), were identified as comprising partial amino acid sequences of a new human hematopoietin receptor polypeptide, HPR1. These amino acids sequences were used to identify a DNA sequence (SEQ ID NO:3) encoding an HPR1 polypeptide having the amino acid sequence shown in SEQ ID NO:4; nucleotides 132 through 2366 of SEQ ID NO:3 encode SEQ ID NO:4, with nucleotides 2367 through 2369 corresponding to a stop codon. The HPR1 coding sequence (nucleotides 132 through 2369 of SEQ ID NO:3) is presented as SEQ ID NO:5. The HPR1 sequences of SEQ ID NOs 3 and 5 were confirmed by three independent PCR amplification experiments from a U937 cDNA library. These HPR1 coding sequences were compared with publicly available preliminary human genomic DNA sequences, and the following chromosome 5 contigs were identified as containing HPR1 coding sequences: AC022265.3, AC008914.3, AC008857.4, and AC016596.4. The human genomic region

5 corresponding to these contigs also includes the gene for gp130, which suggests that gp130 and HPR1 may derive from a common ancestral gene by gene duplication. The approximate positions of the exons containing HPR1 coding sequence in the AC022265.3 contig are shown in the table below, along with their locations relative to SEQ ID NOs 3 and 5; note that the 5' and 3' untranslated regions may extend further along the contig sequence beyond those portions that correspond to SEQ ID NOs 3 and 5, as indicated by the parentheses around the AC022265.3 endpoints in the table. Due to the preliminary sequence and assembly of the contig sequence, the exons within the contig are not always in the right order or orientation with respect to each other, and may contain sequence variations due to inaccurate sequence data or allelic polymorphism.

Corresponding positions of HPR1 gene exons in human contig AC022265.3 and in cDNA sequences:

	Position in AC022265.3	Position in SEQ ID NO:3 / Position in SEQ ID NO:5
Exon 1	(128423)-128559	1-137 / 1-6
Exon 2	134501-134591	138-228 / 7-97
Exon 3	143777-143894	229-346 / 98-215
Exon 4	147256-147437	347-528 / 216-397
Exon 5	51249-51098	529-680 / 398-549
Exon 6	44322-44157	681-846 / 550-715
Exon 7	16473-16394	847-926 / 716-795
Exon 8	30331-30115	927-1143 / 796-1012
Exon 9	178626-178808	1144-1326 / 1013-1195
Exon 10	179879-179980	1327-1428 / 1196-1297
Exon 11	180785-180931	1429-1575 / 1298-1444
Exon 12	183052-183192	1576-1716 / 1445-1585
Exon 13	185997-186090	1717-1810 / 1586-1679
Exon 14	187367-187448	1811-1892 / 1680-1761
Exon 15	189165-(189747)	1893-2480 / 1762-2238

A nucleic acid encoding a polypeptide with a high degree of amino acid similarity (approximately 61% amino acid identity) to human HPR1 was isolated from Mus musculus. The Mus HPR1 amino acid sequence is presented as SEQ ID NO:12, and due to its high level of similarity with human HPR1, is considered to be the murine homologue of human HPR1. PCR amplification of cDNA sequences corresponding to mRNAs encoding murine HPR1 identified a cDNA molecule encoding SEQ ID NO:12; the nucleotide sequence of this murine HPR1 cDNA is presented as SEQ ID NO:28. Nucleotides 1 through 2178 of SEQ ID NO:28 encode SEQ ID NO:12 with publicatides 2170.

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acids 56 through 77 of SEQ ID NO:1 may be a portion of an alternatively spliced exon added following the exon/intron boundary identified between nucleotides 846 and 847 of SEQ ID NO:3 (nucleotides 715 and 716 of SEQ ID NO:5). In an additional potential splice variant, an amino acid sequence ending in the amino acids of SEQ ID NO:10 could be substituted for the amino acids leading up to and including the lysine at position 190 of SEQ ID NO:4. However, such a splice variant would require an additional exon/intron boundary approximately between nucleotides 701 and 702 of SEQ ID NO:3 (nucleotides 570 and 571 of SEQ ID NO:5). In a further potential splice variant, the amino acid sequence of SEQ ID NO:11 could be substituted for amino acids 238 through 266 of SEQ ID NO:4 by replacing exon 7 with an alternative exon encoding the SEQ ID NO:11 amino acids. In this potential variant, 29 amino acids C-terminal to the WSXWS motif and including the N-terminal portion of the most N-terminal fibronectin type III repeat (as shown in Table 1) would be replaced with 15 amino acids, resulting in deletion of a portion of the most N-terminal fibronectin type III repeat, including two highly conserved Trp residues.

Additional variations of HPR1 polypeptides are provided as naturally occurring genomic variants of the HPR1 sequences disclosed herein; such variations may be incorporated into an HPR1 polypeptide or nucleic acid individually or in any combination, or in combination with alternative splice variation as described above. As one example, amino acids 5 through 40 of SEQ ID NO:2 match SEQ ID NO:4, with amino acid 4 of SEQ ID NO:2 likely representing an allelic variation, where the change from the Asn residue position 187 of SEQ ID NO:4 to a Thr residue in SEQ ID NO:2 could be caused by a single change from 'A' to 'C' at position 691 of SEQ ID NO:3 or 560 of SEQ ID NO:5. This variation and others are listed in the table below:

Amino Acid Change	Position in SEQ ID NO:4	Nucleotide Change	Position in SEQ ID NO:3 / Position in SEQ ID NO:5
Thr -> Ala	83	A -> G	378 / 247
Asp -> Asn	168	G-> A	633 / 502
Asn -> Thr	187	A -> C	691 / 560
Ser -> Pro	361	T -> C	1212 / 1081
Ala -> Gly	362	C-> G	1216 / 1085
Ser -> Asn	510	G -> A	1660 / 1529
Asn -> Asp	517	A -> G	1680 / 1549
Arg -> Gly	679	A -> G	2166 / 2035

#### B. Identification of HPR2, a New Member of the Human Hematopoietin Receptor Family

A data set was received from Celera Genomics (Rockville, Maryland) containing a listing of amino acid sequences predicted to be encoded by the human genome. This data set was searched with a BLAST algorithm to identify hematopoietin receptor family polypeptides. Several amino acid sequences, including SEQ ID NO:16, were identified as comprising partial amino acid sequences of a new human hematopoietin receptor polypeptide, HPR2. These amino acids sequences were used to identify a DNA sequence (SEQ ID NO:19) encoding an HPR2 polypeptide having the amino acid sequence shown in SEQ ID NO:21; nucleotides 107 through 1993 of SEQ ID 19 encode SEQ ID

NO:21, with nucleotides 1994 through 1996 corresponding to a stop codon. The HPR2 coding sequence (nucleotides 107 through 1996 of SEQ ID NO:19) is presented as SEQ ID NO:20. The HPR2 sequences of SEQ ID NOs 19 and 20 were confirmed by independent PCR amplification experiments from a human lymph node cDNA library and a CB23 B cell line cDNA library. These PCR amplification experiments also identified two additional splice variants of the HPR2 cDNA sequence referred to as HPR2-ex8-ex9 and HPR2-ex9; the coding sequences for HPR2-ex8-ex9 and HPR2-ex9 are presented as SEQ ID NOs 22 and 24, respectively, and the amino acid sequences they encode are presented as SEQ ID NOs 23 and 25, respectively. The HPR2 cDNA sequences of SEQ ID NOs 19, 20, and the HPR2-ex8-ex9 cDNA of SEQ ID NO:22 were present in both the lymph node and CB23 cDNA libraries, while the HPR2-ex9 cDNA of SEQ ID NO:24 was only present in the lymph node library.

These HPR2 coding sequences were compared with publicly available preliminary human genomic DNA sequences, and the following chromosome 1 contigs were identified as containing HPR2 coding sequences: GenBank accession numbers AL109843 (1p31.2-32.1) and AL389925. The human genomic region corresponding to the AL389925 contig also includes the gene for IL-12RB2, which suggests that IL-12RB2 and HPR2 may derive from a common ancestral gene by gene duplication. The approximate positions of the exons containing HPR2 coding sequence in the AL109843 and AL389925 contigs are shown in the table below, along with their locations relative to SEQ ID NOs 19, 20, 22, and 24; note that the 5' and 3' untranslated regions may extend further along the contig sequence beyond those portions that correspond to SEQ ID NOs 19, 20, 22, and 24, as indicated by the parentheses around the AL109843 and AL389925 endpoints in the table. Due to the preliminary nature of the sequence data and assembly of the contig sequence, the exons within the genomic contigs may contain sequence variations due to inaccurate sequence data or allelic polymorphism.

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Corresponding positions of HPR2 gene exons in human genomic contigs AL109843 and AL389925 and in HPR2 coding sequences:

	Position in AL109843	Position in SEQ ID NO:19 / 20 / 22 / 24
Exon 1	(34088)-34164	1-77 / (5' UTR, not in SEQ ID NOs 20, 22, and 24)
Exon 2	35715-35813	78-176 / 1-70 / 1-70 / 1-70
Exon 3	36965-37261	177-473 / 71-367 / 71-367 / 71-367
Exon 4	50459-50582	474-597 / 368-491 / 368-491 / 368-491
Exon 5	68360-68520	598-758 / 492-652 / 492-652 / 492-652
Exon 6	74533-74678	759-904 / 653-798 / 653-798 / 653-798
Exon 7	87197-87353	905-1061 / 799-955 / 799-955 / 799-955
Exon 8	104336-104425	1062-1151 / 956-1045 / (not present) / 956-1045
Exon 9	107802-107904	1152-1254 / 1046-1148 / (not present) / (not present)
	Position in AL389925	Position in SEQ ID NO:19 / 20 / 22 / 24
Exon 10	8847-8937	1255-1345 / 1149-1239 / 'G'-957-1047 / 1046-1071
Exon 11	11488-(12972)	1346-2830 / 1240-1890 / 1048-1698 / (not present)

In the HPR3-ex9 splice variant, note that the absence of the exon 9 sequence (103 nucleotides) changes the reading frame towards the 3' end of the coding sequence for the HPR2-ex9 form (SEQ ID NO:24)

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relative to that of the HPR2 coding sequence of SEQ ID NO:20, leading to a different amino acid sequence in the HPR2-ex9 C-terminal portion and a stop codon after amino acid 356 (compared to 629 amino acids in HPR2). For the HPR2-ex8-ex9 form, the splice is made at a slightly different exon 10 splice acceptor site than for the HPR2 form, so that an extra 'G' residue is included at the start of exon 10 in the HPR2-ex8-ex9 form, restoring the reading frame to be the same as in the 3' end of the HPR2 sequence. The C-terminal 248 amino acids of HPR2-ex8-ex9 form are therefore the same as the C-terminal 248 amino acids of HPR2 form, and although the coding sequence of the HPR2-ex8-ex9 form is missing both exons 8 and 9 (except for the last 'G' residue of exon 9), the resulting HPR2-ex8-ex9 form polypeptide is longer (565 amino acids) than the HPR2-ex9 form polypeptide (356 amino acids).

Several splice variations of the HPR2 sequences have been identified in human genomic sequences and are included within the scope of the invention. For example, amino acids 118 through 215 of SEQ ID NO:16 match the amino acid sequence of HPR2 presented in SEQ ID NO:21, while amino acids 1 through 117 of SEQ ID NO:16 may correspond to an alternatively spliced exon added upstream of exon 3 (i.e. at the exon/intron boundary identified between nucleotides 176 and 177 of SEQ ID NO:19). Amino acids 216 through 245 of SEQ ID NO:16 may correspond to an additional alternatively spliced exon added between exon 3 and exon 4 (i.e. at the exon/intron boundary identified between nucleotides 473 and 474 of SEQ ID NO:19). Amino acids 340 through 344 of SEQ ID NO:16 may correspond to an alternatively spliced exon added downstream of exon 5 (i.e. at the exon/intron boundary identified between nucleotides 758 and 759 of SEQ ID NO:19). In a further potential splice variant, an alternative exon or exons encoding the amino acid sequence of SEQ ID NO:17 could be substituted for exon 6, resulting in the replacement of amino acids 217 through 267 of SEQ ID NO:21 with the SEQ ID NO:17 amino acids. In this potential variant, 51 amino acids N-terminal to the WSXWS motif, including the proline-rich region (as shown in Table 1) between the two cytokine receptor subdomains, would be replaced with 39 amino acids, resulting in deletion of a portion of the more C-terminal cytokine receptor subdomain which includes a highly conserved Trp residue. In an additional potential splice variant, an alternative exon could be added downstream of exon 4 (i.e. at the exon/intron boundary identified between nucleotides 597 and 598 of SEQ ID NO:19) so that an amino acid sequence starting in the amino acids of SEQ ID NO:18 could be substituted for amino acids following and including the serine at position 164 of SEQ ID NO:21. Multiple splice variations as described above can be included in a single splice variant, for example, replacing exon 6 with an alternative exon or exons encoding the amino acid sequence of SEQ ID NO:17, and also deleting exons 8 and/or 9 as described above.

Additional variations of HPR2 polypeptides are provided as naturally occurring genomic variants of the HPR2 sequences disclosed herein; such variations may be incorporated into an HPR2 polypeptide or nucleic acid individually or in any combination, or in combination with alternative splice variation as described above. As one example, a change from the Leu residue position 310 of SEQ ID NO:21 to a Pro residue could be caused by a single change from "T" to 'C' at position 1035 of SEQ ID NO:19. This variation and another are listed in the table below:

Amino Acid Change	Position in SEQ ID NO:21	Nucleotide Change	Position in SEQ ID NO:19
Leu->Pro	310	T->C	1035
(not applicable)	(not applicable)	A->G	2172 (3' UTR)

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A nucleic acid encoding a polypeptide with a high degree of amino acid similarity (approximately 69% amino acid identity) to human HPR2 was isolated from Mus musculus. The Mus HPR2 amino acid sequence is presented as SEQ ID NO:27, and due to its high level of similarity with human HPR2, is considered to be the murine homologue of human HPR2. PCR amplification of cDNA sequences corresponding to mRNAs encoding murine HPR2 identified a cDNA molecule encoding SEQ ID NO:27; the nucleotide sequence of this murine HPR2 cDNA is presented as SEQ ID NO:29. Nucleotides 1 through 1932 of SEQ ID NO:29 encode SEQ ID NO:27, with nucleotides 1933-1935 corresponding to a stop codon. The murine HPR2 amino acid sequence of SEO ID NO:27 appears to have a 20-amino acid insertion at amino acids 297 through 316 of SEQ ID NO:27 relative to human HPR2 of SEQ ID NO:21, based on an alignment of the human and murine polypeptide sequences; this insertion is identical to amino acids 317 through 336. Given the number of alternatively spliced forms identified for human HPR2, it is possible that this insertion in murine HPR2 relative to the human HPR2 of SEQ ID NO:21 is the result of alternative splicing. One embodiment of the invention is a form of murine HPR2 in which one of these repeated WQPWS-containing motifs has been deleted; that is, polypeptides in which the amino acid sequence ending with amino acid 296 of SEQ ID NO:27 is contiguous with the amino acid sequence beginning with amino acid 317 of SEQ ID NO:27, or polypeptides in which the amino acid sequence ending with amino acid 316 of SEQ ID NO:27 is contiguous with the amino acid sequence beginning with amino acid 337 of SEQ ID NO:27.

#### C. Comparison of HPR1 and HPR2 to Other Hematopoietin Receptor Polypeptides.

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The amino acid sequences of human HPR1 (SEQ ID NO:4), murine HPR1 (SEQ ID NO:12), and human HPR2 (SEQ ID NO:21) were compared with the amino acid sequences of these other hematopoietin receptor family members - LIF-R, the interleukin 12 beta 2 receptor chain (IL-12RB2), gp130, and GCSFR (SEQ ID NO:6 - SEQ ID NO:9, respectively) - using the GCG "pretty" multiple sequence alignment program, with amino acid similarity scoring matrix = blosum62, gap creation penalty = 8, and gap extension penalty = 2. Alignments of these sequences are shown in Table 1, and include consensus residues which are identical among at least three of the amino acid sequences in the alignment. The capitalized residues in the alignment are those which match the consensus residues. The numbering of amino acid residues in Table 1 corresponds to the position of those residues in the HPR1 amino acid sequence (SEQ ID NO:4). Note that only a portion of the HPR2 amino acid sequence is shown in Table 1, as HPR2 does not contain fibronectin type III repeats in its extracellular domain. HPR1 and HPR2 sequences corresponding to the intracellular Box 1 and Box 2 motifs are shown in Table 2. Sequences of eleven amino acids similar to the Box 1 or 2 motif of other hematopoietin receptors were identified for HPR1 and HPR2, and placed into a column with these motif sequences (with no gaps introduced). Similarly, HPR2 sequences corresponding to the

5 intracellular Box 3 motif are shown in Table 3. Sequences of fourteen amino acids similar to the Box 3 motif of other hematopoietin receptors were identified for HPR2, and placed into a column with these motif sequences (with no gaps introduced). The numbering of each sequence on Tables 2 and 3 corresponds to their position in the complete amino acid sequence for that HPR polypeptide. The consensus residues are those that are present in three or more (for Table 2) or two or more (for Table 3) sequences at that position in the motif.

Amino acid substitutions and other alterations (deletions, insertions, etc.) to HPR1 and HPR2 amino acid sequences (for example, SEQ ID NOs 4, 12, and 21) are predicted to be more likely to alter or disrupt HPR1 or HPR2 polypeptide activities if they result in changes to the capitalized residues of the amino acid sequences as shown in Tables 1, 2, and 3, and particularly if those changes do not substitute an amino acid of similar structure (such as substitution of any one of the aliphatic residues -Ala, Gly, Leu, Ile, or Val - for another aliphatic residue), or a residue present in other hematopoietin receptor polypeptides at that conserved position. Conversely, if a change is made to an HPR1 or HPR2 amino acid sequence resulting in substitution of the residue at that position in the alignment from one of the other Table 1, 2, or 3 hematopoietin receptor polypeptide sequences, it is less likely that such an alteration will affect the function of the altered HPR1 or HPR2 polypeptide. For example, the consensus residue at position 42 in Table 1 is serine, and one of the hematopoietin receptors (LIF-R) has an asparagine at that position. Substitution of asparagine or the chemically similar glutamine for serine at that position is considered to be less likely to alter the function of the polypeptide than substitution of tryptophan or tyrosine etc. Embodiments of the invention include HPR1 and HPR2 polypeptides and fragments of HPR1 and HPR2 polypeptides, comprising altered amino acid sequences. Altered HPR1 or HPR2 polypeptide sequences share at least 30%, or more preferably at least 40%, or more preferably at least 50%, or more preferably at least 55%, or more preferably at least 60%, or more preferably at least 65%, or more preferably at least 70%, or more preferably at least 75%, or more preferably at least 80%, or more preferably at least 85%, or more preferably at least 90%, or more preferably at least 95%, or more preferably at least 97.5%, or more preferably at least 99%, or most preferably at least 99.5% amino acid identity with one or more of the hematopoietin receptor amino acid sequences shown in Tables 1, 2, and 3.

Table 1: Alignment of HPR1 and HPR2 extracellular domains with those of other hematopoietin receptors

35 C: conserved cysteine

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: proline-rich 'linker' between cytokine receptor subdomains

: WSXWS motif

: fibronectin type III repeats

	SEQ ID NO:	35	78
Hs HPR1	4	PakPeNiSCV yYyr.KNLTC TWS	PGkETsyTqYTv KrtyafGekh
Mus HPR1	12	PtKPeNiSCV fYfd.rNLTC TWr	PekETndTSYiv tltySyGK
gp130	8	PekPkNLSCi vneg.KkmrC eWd	gGrETHLeTnfTL KsewathKfa
GCSFR	9	PaiPhNLSCl mnlttssLiC qWe	PGpETHLpTSfTL KsfkSrGnCq
Hs HPR2	21	PdiPdevtCV iYeysgNmTC TWn	aGklTyidTkYvv hvKsl

LIF-R consensus	<u>7</u>	DOWDWITT CO:				
	_	Lededwrgc1	qkgeqgtva <b>C</b>	TWerGrdTHL	yTeYTL	qlSgpKnl
consensus	Ö	PdtPqqLn <b>C</b> e	th.dlKeiiC	sWnPGrvTaL	vgpraTSYTL	vesfS.GKyv
		P-KP-NLSCV	-YKNLTC	TW-PG-ETHL	TSYTL	KS-GK
			. <del> </del>			<del></del>
<del></del>		79				126
Hs HPR1	4		on so affi	PRitipdN	Whi oldow Edia	
	12	anculasis	enraststin	PRILIpun	TCTEVEALNG	dGVIKSIIIILY
Mus HPR1		snys	dnaceasysi	PRscamppdi	csvevQAqNg	agkvksalty
gp130	8			stvyfvN		
GCSFR	9			PRkhlllyqN		
Hs HPR2	21	eteeeggylt	ssyini	stdsląggkk	YlVWVQAaNA	LGmeeSkqLq
IL-12RB2	7			tpespesN		
LIF-R	6			Pnqei		
consensus	_ <u>~</u>				Y-VWVQAENA	
COLLECTIONS					T 444 0 521111111	70 b b 7
		127				172
			Dol wonter	<del></del>	101 7711 7 7	
Hs HPR1	4	wrLenlaktE		lgikrm		apvssdLKyt
Mus HPR1	12	whLisIaKtE		cnrm		ktrgfpLvCm
gp130	8	fdpvykVKPn	PPhnlsVins	eelssi	<u>lkLtWtnPsi</u>	ksv.liLKyn
GCSFR	9	1dpmDvVK1E	PPmlrtmdPs	peaappqagc	lQLcW.ePwq	pglhIngKCE
Hs HPR2	21			natvokti		
IL-12RB2	<del></del> -			kasvsrct		
LIF-R	6			nsta		nfakInf1CE
consensus		L-DIVKPE	DDTDVVDT			I-LKCE
Consensus		DIAKER			-Оп-м	
			*** <u></u>			
		173			·	221
Hs HPR1	4			rkdknqtynL		
Mus HPR1	12			ckqvcnL		
gp130	8	iqYRTkda.s	tWsqip.ped	tastrssftv	qdLkPFTEYV	FrIRCmkEdg
GCSFR	9	LRhkpgrgea	sWalVgp	lplealqyeL	cGLlPaTaYt	lOIRCirwpl
Hs HPR2	21	mRYkattnqt	.WnvkeFd	tnftvvggse	fvLePnikYV	FQvRCq.Etg
	7			vickakarnal	TOTKELLIFAE	MOISSKINIV
IL-12RB2	7			vtkakgrhdL		FQIssklhly
IL-12RB2 LIF-R	6	ieikksNSvq	eqrnVti.kg	venssylvaL	dkLnPyTlYt	FrIRCstEtf
IL-12RB2		ieikksNSvq	eqrnVti.kg		dkLnPyTlYt	FrIRCstEtf
IL-12RB2 LIF-R		ieikksNSvq	eqrnVti.kg	venssylvaL	dkLnPyTlYt	FrIRCstEtf
IL-12RB2 LIF-R		ieikksNSvq LRYRT-NS	eqrnVti.kg	venssylvaL	dkLnPyTlYt	FrIRCstEtf FQIRCE
IL-12RB2 LIF-R consensus	6	ieikksNSvq LRYRT-NS	eqrnVti.kg -WV-FN	venssylvaL	dkLnPyT1Yt -GL-PFTEYV	FrIRCstEtf FQIRCE
IL-12RB2 LIF-R Consensus Hs HPR1	4	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE	eqrnVti.kg -WV-FN kmgmTeEEaP	venssylvaL	dkLnPyTlYt -GL-PFTEYV lWRvLkP	Frircstetf FOIRCE 261 aeadGrRpVr
LL-12RB2 LIF-R Consensus Hs HPR1 Mus HPR1	6 4 12	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE	eqrnVti.kg -WV-FN kmgmTeEEaP etrvTmEEvP	venssylvaLL cgLe hvLD	dkLnPyTlYt -GL-PFTEYV  IWRvLkP IWRiLeP	FrIRCstEtf FOIRCE 261 aeadGrRpVr admnGdRkVr
IL-12RB2 LIF-R Consensus Hs HPR1	6 4 12 8	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE	eqrnVti.kg -WV-FN kmgmTeEEaP etrvTmEEvP	venssylvaLL cgLe hvLD	dkLnPyTlYt -GL-PFTEYV  IWRvLkP IWRiLeP	FrIRCstEtf FOIRCE 261 aeadGrRpVr admnGdRkVr
LL-12RB2 LIF-R Consensus Hs HPR1 Mus HPR1	6 4 12	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE	eqrnVti.kg -WV-FN kmgmTeEEaP etrvTmEEvP asgiTyEdrP	cgLe hvLD skaps	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP	FrIRCstEtf FOIRCE 261 aeadGrRpVr admnGdRkVr shtqGyRtVq
LL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130	6 4 12 8 9	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP	cgLe hvLD skaps tvrLD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP	FrIRCstEtf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 gp130 GCSFR HS HPR2	4 12 8 9 21	ieikksNSvq LRYRT-NS 222 K.fWSDWSQE r.YWSkWSKE KGYWSDWSPE pGhWSDWSPs KrYWqpWSsl	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP le1rTtEraP ffhkTpEtvP	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhD	dkLnPyTlYt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~	FrIRCstEtf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 gp130 GCSFR HS HPR2 IL-12RB2	4 12 8 9 21 7	ieikksNSvq LRYRT-NS 222 K.fWSDWSQE r.YWSkWSkE KGYWSDWSPE pGhWSDWSPS KrYWQPWSsl KGSWSDWSes	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP le1rTtEraP ffhkTpEtvP lraqTpEEeP	cgLe hvLD skaps tvrLD qvtskafqhDtgmlD	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD.	FrIRCstEtf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqis
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R	4 12 8 9 21	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP le1rTtEraP ffhkTpEtvP 1raqTpEEeP kqh1TtEasP	cgLe hvLD skaps tvrLD qvtskafqhDtgmlD skgpD	dkLnPyT1Yt -GL-PFTEYV  lwRvLkP lwRiLeP fwykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. Twrew	FrIRCstEtf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknli
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 gp130 GCSFR HS HPR2 IL-12RB2	4 12 8 9 21 7	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP le1rTtEraP ffhkTpEtvP 1raqTpEEeP kqh1TtEasP	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TW-~-R-LDP	FrIRCstEtf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R	4 12 8 9 21 7	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP le1rTtEraP ffhkTpEtvP 1raqTpEEeP kqh1TtEasP	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TW-~-R-LDP	FrIRCstEtf FQIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS	4 12 8 9 21 7 6	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpD	dkLnPyTlYt -GL-PFTEYV  lwRvLkP lwRiLeP fwykiDP TwwrqRLDP Twns~~~~ vWymkRhiD. Twrew TW-~-R-LDP	FrIRCstEtf FQIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1	4 12 8 9 21 7	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD	dkLnPyTlYt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWms~~~~ vWymkRhiD. TWrew TWR-LDP	FrIRCstEtf FQIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-  310 qlelhLgges
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 MUS HPR1	4 12 8 9 21 7 6	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSPE PGHWSDWSPS KrYWqPWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn.	dkLnPyT1Yt -GL-PFTEYV  lwRvLkP lwRiLeP fwykiDP TwwrqRdDP Twms~~~~ vWymkRhiD. Twrew Tw-~R-LDP	FrIRCstEtf FQIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1	6 12 8 9 21 7 6	ieikksNSvq LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSPE PGHWSDWSPS KrYWqPWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD	dkLnPyT1Yt -GL-PFTEYV  lwRvLkP lwRiLeP fwykiDP TwwrqRdDP Twms~~~~ vWymkRhiD. Twrew Tw-~R-LDP	FrIRCstEtf FQIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 MUS HPR1	6 12 8 9 21 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LvWktLPpfE	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsqqaga	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TW-~-R-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT. ilplcNtTel	FrIRCstEtf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea
LL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GD130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 MUS HPR1 GD130 GCSFR	6 12 8 9 21 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LvWktLPpfE	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsqqaga	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TW-~-R-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT. ilplcNtTel	FrIRCstEtf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 MUS HPR1 GP130 GCSFR IL-12RB2	6 12 8 9 21 7 6 4 12 8 9 7	ieikksNSvq LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSbwSnk KGSWSDWS-E  262 L1WKkargap L1WKkargap LVWKtLPpfE LfWKpvPleE	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka	dkLnPyTlYt -GL-PFTEYV  IWRvLkP 1WRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts	FrIRCstEtf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 MUS HPR1 GP130 GCSFR IL-12RB2 LIF-R	6 12 8 9 21 7 6	ieikksNSvq LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGSWSDWS-E  262 L1WKkargap L1WKkargap L1WKktLPpfE LfWKpVPleE LfWKpLSVSE	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV AnGKILSYnV	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTgs	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh	Frircstetf FOIRCE  261 aeadGrRpVr admmGdRkVr shtqGyRtVq,RtVq,ysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdknd
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 MUS HPR1 GP130 GCSFR IL-12RB2	6 12 8 9 21 7 6 4 12 8 9 7	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262  LIWKkargap LIWKkargap LVWKtLPpfE LfWKpVPleE LfWKpLPinE LfWKpLPinE	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AngKILdYeV dsGrIqGYvV ArGKILhYqV AngKILsYnV A-GKILGY-N	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsqaga tlqeltggka scssdeeTqs	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~ vWymkRhiD. TWrew TWR-LDP  LTEINNITTq qnytvNaT ilplcNtTel mTqnitghts LSEipd.pqh LTE-N-T-	Frircstetf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVq,RtVq,ysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtga kaeirLdknd
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 MUS HPR1 GP130 GCSFR IL-12RB2 LIF-R	6 12 8 9 21 7 6 4 12 8 9 7	ieikksNSvq LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 L1WKkargap L1WKkargap LVWKtLPpfE LfWKpVPleE LfWKpLSVSE iyWKpLPinE L-WK-LPE	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AngKILdYeV dsGrIqGYvV ArGKILhYqV AngKILsYnV A-GKILGY-N	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsqaga tlqeltggka scssdeeTqs	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~ vWymkRhiD. TWrew TWR-LDP  LTEINNITTq qnytvNaT ilplcNtTel mTqnitghts LSEipd.pqh LTE-N-T-	Frircstetf FOIRCE  261 aeadGrRpVr admmGdRkVr shtqGyRtVq,RtVq,ysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR CONSENSUS  LIF-R CONSENSUS	4 12 8 9 21 7 6 4 12 8 9 9 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262  L1WKkargap L1WKkargap LVWKtLPpfE LfWKpLPPiee LfWKpLPine LfWKpLP-E	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AngKILGYeV dsGriqGyeV ArgKILhyqV ArgKILhyqV ArgKILBGY-V	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsqaga tlqeltggka scsdeeTqsT	dkLnPyT1Yt -GL-PFTEYV  1WRvLkP 1WRiLeP fWykiDP TWwrqRqLDP TWns~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTEN-T	Frircstetf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqgkVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdknd
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR IL-12RB2 LIF-R CONSENSUS  HS HPR1 CONSENSUS  HS HPR1	4 12 8 9 21 7 6 4 12 8 9 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262  LlWKkargap LlWKkargap LVWKtLPpfE LfWKpLPinE LfWKpLPinE LfWKpLPinE L-WK-LP-E	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AngKILdYeV dsGriqGYeV ArgKiLhyqV ArgKiLhyqV AngKILsYnV A-GKILGY-V lGKSpvatLr	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgaga tlqeltggka scssdeeTqsT	dkLnPyT1Yt -GL-PFTEYV  lwRvLkP lwRiLeP fwykiDP TWwrqRqLDP TWns~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T	Prircstetf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GD130 GCSFR IL-12RB2 LIF-R CONSENSUS  HS HPR1 GD130 GCSFR IL-12RB2 LIF-R CONSENSUS	4 12 8 9 21 7 6 4 12 8 9 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262  LlWKkargap LlWKkargap LVWKtLPpfE LfWKpVPleE LfWKpLPinE L-WK-LPE  311  fwVSmisyNS hsVSVtsfNS	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILSYnV A-GKILSYNV A-GKILGY-V  1GKSpvatlr 1GKSqeTilr	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqs	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWmrax vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTEN-T	Prircstetf FOIRCE  261 aeadGrRpVr admnGdrkVr shtqGyrtVqrkVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdknd
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GCSFR IL-12RB2 LIF-R CONSENSUS  HS HPR1 GCSFR IL-12RB2 LIF-R CONSENSUS	6 4 12 8 9 21 7 6 4 12 8 9 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KryWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262  LlWKkargap LlWKkargap LVWKtLPpfE LfWKpVPleE LfWKnLsvsE iyWKpLPinE L-WK-LPE  311  fwVSmisyNS hsVSVtsfNS ylatltvrNl	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILSYnV A-GKILSYnV LGKSdaavLt	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgaga tlqeltggka scssdeeTqs	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWwrqRqLDP TWms~~~~ vWymkRhiD. TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT. ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T cievmqAcva yiksmqAyia pvmdlkAfpk	Prircstetf FOIRCE  261 aeadGrRpVr admnGdrkVr shtqGyrtVqrvqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVewt
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS  HS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS	4 12 8 9 21 7 6 4 12 8 9 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262  LlWKkargap LlWKkargap LlWKkargap LlWKkLPpfE LfWKpLPinE LfWKpLPinE L-WK-LPE  311  fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILSYnV A-GKILGY-V  lGKSpvatLr lGKSqeTiLr vGKSdaavLt aGtSrPTpv.	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgaga tlqeltggka scssdeeTqsT IPaiqEksfq IPacdfqath .vfsEsrgp	dkLnPyTlYt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWwrqRqLDP TWms~~~~ vWymkRhiD. TWrew TWR-LDP  LTEINNITTq qnytvNaT. ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T cievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar	Prircstetf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt dphsLwVgWe
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS  HS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS	6 4 12 8 9 21 7 6 4 12 8 9 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262  LlWKkargap LlWKkargap LlWKkargap LlWKkLPpfE LfWKpLPinE LfWKpLPinE L-WK-LPE  311  fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILSYnV A-GKILGY-V  lGKSpvatLr lGKSqeTiLr vGKSdaavLt aGtSrPTpv.	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgaga tlqeltggka scssdeeTqs	dkLnPyTlYt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWwrqRqLDP TWms~~~~ vWymkRhiD. TWrew TWR-LDP  LTEINNITTq qnytvNaT. ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T cievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar	Prircstetf FOIRCE  261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt dphsLwVgWe
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS  HS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS	4 12 8 9 21 7 6 4 12 8 9 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWS-E .wkWSkWSnk KGYWSDWS-E 262 LlWKkargap LlWKkargap LlWKkargap LVWKtLPpfE LfWKpLPinE LfWKpLPinE L-WK-LPE 311 fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS waVaVsAaNS	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILSYnV A-GKILGY-V  1GKSpvatLr 1GKSqeTiLr vGKSdaavLt aGtSrPTpv. kGsSlPTrin	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgaga tlqeltggka scssdeeTqs	dkLnPyTlYt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWwrqRqLDP TWms~~~~ vWymkRhiD. TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT. ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T cievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar aprqvsAnse	Frircstetf FOIRCE  261 aeadGrRpVr admnGdrkVr shtqGyrtVqrtVqysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt dphsLwVgWe gmdnilVtWQ
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS  HS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS	6 4 12 8 9 21 7 6 4 12 8 9 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262  LlWKkargap LlWKkargap LlWKkargap LlWKkargap LVWKtLPpfE LfWKpLPinE LfWKpLPinE L-WK-LPE  311 fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS waVaVsAaNS yiiSVvAkNS	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILSYNV A-GKILGY-V  lGKSpvatlr lGKSpvatlr vGKSdaavLt aGtSrPTpv. kGsSlPTrin vGsSpPskia	venssylval  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn qyfaensTn tltrwkshl swrpsgaga tlqeltggka scssdeeTqsT- IPaiqEksfq IPacdfqath .vfsEsrgp ImnlcEagll smeipnddl.	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT. ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T CievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar aprqvsAnsekieqvvg	Prircstetf FOIRCE  261 aeadGrRpVr admmGdRkVr shtqGyRtVq,RtVq,ysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt dphsLwVgWe gmdnilVtWQ mgkgilltWh
IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR HS HPR2 IL-12RB2 LIF-R CONSENSUS  HS HPR1 MUS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS  HS HPR1 GP130 GCSFR IL-12RB2 LIF-R CONSENSUS	6 4 12 8 9 21 7 6 4 12 8 9 7 6	ieikksNSvq LRYRT-NS  222  K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWS-E .wkWSkWSnk KGYWSDWS-E 262 LlWKkargap LlWKkargap LlWKkargap LVWKtLPpfE LfWKpLPinE LfWKpLPinE L-WK-LPE 311 fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS waVaVsAaNS	eqrnVti.kg -WV-FN  kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILSYNV A-GKILGY-V  lGKSpvatlr lGKSpvatlr vGKSdaavLt aGtSrPTpv. kGsSlPTrin vGsSpPskia -GKS-PT-L-	venssylval  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgaga tlqeltggka scssdeeTqsT-  IPaiqEksfq IPdvhEktfq IPacdfqath .vfsEsrgp ImnlcEagll smeipnddl. IPE	dkLnPyT1Yt -GL-PFTEYV  IWRvLkP IWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT. ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T CievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar aprqvsAnsekieqvvgA	Prircstetf FOIRCE  261 aeadGrRpVr admmGdRkVr shtqGyRtVq,RtVq,ysrqqisssdgknliG-R-V-  310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt dphsLwVgWe gmdnilVtWQ mgkgilltWh

	f	360 404	1
Hs HPR1	4	ssaldVn twmIEWfpdv d.SePttlsW e.svSqaTnw TIqqDkLKPE	7
Mus HPR1	12	ssipaVd twivEWlpea amSkfpalsW e.svSqvTnw TleqDkLKPF	
gp130	8	tPresVk kYilEWcvls dka.PcitdW q.qedgtvhr TylrgNLaes	<u>-</u> -
GCSFR	9	pPnpwpq gYVIEWglgp psasnsnktW rmeqngratg fllkeNirPE	<u> </u>
IL-12RB2	7	pPrkdpsaVq eYVvEWrelh pggdtqvpln wlrsrpynvs aliseNiKsy	
LIF-R	6	ydpnmtc dYVIkWc.ns srSePclmdW rkvpSnsTet vIesDefrPc	<u>J</u>
consensus		PVYVIEWS-PWS-T TIDNLKPE	7
			_
		405 454	1
Hs HPR1	4	wCYNIsVYPm lhDkvGePyS IqAYaKEgvP SeGPEtkVEn IGvktvtItW	V_
Mus HPR1	12	tCYNIsVYPv lghrvGePyS IqAYaKEgtP lkGPEtrVEn IGlrtAtItW	1
gp130	8	kCYlItVtPv yaDgpGsPeS IkAYlKqapP SkGPtvrtkk vGKneAvleW	
GCSFR	9	qlYeIiVtPl yqDtmGpsqh vyAYsqEmAP ShaPElhlkh IGKtwAqleV	1
IL-12RB2	7	iCYeIrVYal sgDq.GgcsS IlgnsKhkAP lsGPhinait eeKgsilIsV	V.
LIF-R	6	irYNfflYgc rnqgyqllrS migYieElAP ivaPnftVEd tsadsilvkW	٧
consensus	l	-CYNI-VYPDG-P-S I-AY-KE-AP S-GPEVE- IGKA-I-V	1
			_
l 	<u> </u>	455 489	
Hs HPR1	4	keIPksErkG iicNYTIFYq aeGGkgfSKt	<u>i</u> _
Mus HPR1	12	keIPksarnG FinNYTvFYq aeGGkelSKt	_
gp130	8	dqlPVdvqnG FirNYTIFYr tiiGnetavnVdSSh	<u>.1</u>
GCSFR	9	vpePpelgks plthYTIFwt nagngsfSai	
IL-12RB2	7	nsIPVqEqmG cllhYrIywk erdsnsqpqlceipyrvSc	
LIF-R	6	edIPVeElrG FlrgY.lFYf gkGerdtSKm rvlesgrsdi kVknitdiSc	_
consensus		IPV-EG FNYTIFYGGSKVNSS-	
	<u> </u>		=
	L	490 539	
Hs HPR1	4	lqygLeSLkr kTSYiVqvMA sTsAGGTNGt sinFkTLsfS VfEIilItsI	
Mus HPR1	12	lqcdLeSLtr rTSYtVwvMA sTrAGGTNGv rinFkTLsiS VfEvvlltsI	
gp130	8	teytLsSLts dTlYmVrmaA yTdeGGkdGp eftFtTpkfa qgEIeaIvvr	
GCSFR	9	rgfvLhgLep aslYhihlMA asqAGaTNst vltlmTLtpe gsElhiIlgI	
IL-12RB2	7	nshpinSLqp rvtYvlwmtA lTaAGesshg nerefcLqgk anwmafvaps	
LIF-R	6	ktlriadLqq kTSYhlvlrA yTdqGvqpek smyvvTkenS VqlIiaIlir	_
consensus	ļ	L-SLTSY-VMA -T-AGGTNGF-TLS V-EIII	

Table 2: Box 1 and Box 2 motifs in the intracellular domains of HPR1, HPR2, and other hematopoietin receptors

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	SEQ ID NO	Box 1 Motif	Box 2 Motif
Hs HPR1	4	563-thlcWPtVPNP-573	631-eifTdEArtgq-641
Mus HPR1	12	517-tplccPDVPNP-527	582-VvlTEEAgKgq-592
HPR2	21	393-pkwlyeDi <b>P</b> Nm-403	430-VdpmiteiKei-440
LIF-R	6	866-KetfyPDi <b>P</b> NP-876	910-VleTrsAfpKi-920
gp130	8	648-KkhiWPnV <b>P</b> dP-658	693-VveiEandKKp-703
GCSFR	9	655-KnplWPsV <b>P</b> dP-665	696-ltvlEEdeKKp-706
consensus		KWPDV <b>P</b> NP	VTEEA-KK-

# 10 Table 3: Box 3 motifs in the intracellular domains of HPR2 and other hematopoietin receptors

	SEQ ID NO	Box 3 Motif
HPR2 (first occurrence)	21	478-PdLntGYKPQisnf-491
HPR2 (second occurrence)	21	605-lpsint <b>Y</b> fP <b>Q</b> niLe-618
LIF-R	6	995-PVggaGYKPQmhLp-1008
gp130	8	693-tVvhsG <b>Y</b> rh <b>Q</b> vpsv-774
GCSFR	9	696-PtLvqt <b>Y</b> vl <b>Q</b> gdpr-734
consensus residues		PVL~-G <b>Y</b> KP <b>Q</b> L-

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#### **EXAMPLE 2: Monoclonal Antibodies That Bind Polypeptides of the Invention**

This example illustrates a method for preparing monoclonal antibodies that bind HPR1 or HPR2 polypeptides. Suitable immunogens that may be employed in generating such antibodies include, but are not limited to, purified HPR1 or HPR2 polypeptide or an immunogenic fragment thereof. Purified HPR1 or HPR2 polypeptide can be used to generate monoclonal antibodies immunoreactive therewith, using conventional techniques such as those described in U.S. Patent 4,411,993. Briefly, mice are immunized with HPR1 or HPR2 polypeptide immunogen emulsified in complete Freund's adjuvant, and injected in amounts ranging from about 10 to about 100 micrograms subcutaneously or intraperitoneally. Ten to twelve days later, the immunized animals are boosted with additional HPR1 or HPR2 polypeptide emulsified in incomplete Freund's adjuvant. Mice are periodically boosted thereafter on a weekly to bi-weekly immunization schedule. Serum samples are periodically taken by retro-orbital bleeding or tail-tip excision to test for anti-HPR1 or anti-HPR2 antibodies by dot blot assay, ELISA (Enzyme-Linked Immunosorbent Assay), or inhibition of binding of HPR1 or HPR2 polypeptide to an HPR1 and/or HPR2 binding partner.

Following detection of an appropriate antibody titer, positive animals are provided one last intravenous injection of HPR1 or HPR2 polypeptide in saline. Three to four days later, the animals are sacrificed, spleen cells harvested, and spleen cells are fused to a murine myeloma cell line, e.g., NS1 or preferably P3x63Ag8.653 (ATCC CRL 1580). Fusions generate hybridoma cells, which are plated in multiple microtiter plates in a HAT (hypoxanthine, aminopterin and thymidine) selective medium to inhibit proliferation of non-fused cells, myeloma hybrids, and spleen cell hybrids.

The hybridoma cells are screened by ELISA for reactivity against purified HPR1 or HPR2 polypeptide by adaptations of the techniques disclosed in Engvall et al., (Immunochem. 8:871, 1971) and in U.S. Patent 4,703,004. A preferred screening technique is the antibody capture technique described in Beckmann et al., (J. Immunol. 144:4212, 1990). Positive hybridoma cells can be injected intraperitoneally into syngeneic BALB/c mice to produce ascites containing high concentrations of anti-HPR1 or anti-HPR2 monoclonal antibodies. Alternatively, hybridoma cells can be grown in vitro in flasks or roller bottles by various techniques. Monoclonal antibodies produced in mouse ascites can be purified by ammonium sulfate precipitation, followed by gel exclusion chromatography. Alternatively, affinity chromatography based upon binding of antibody to Polypeptide A or Polypeptide G can also be used, as can affinity chromatography based upon binding to HPR1 or HPR2 polypeptide.

#### **EXAMPLE 3**

#### Antisense Inhibition of HPR1 and/or HPR2 Expression

In accordance with the present invention, a series of oligonucleotides are designed to target different regions of HPR1 and/or HPR2 human or murine mRNA molecules, using the nucleotide sequences of SEQ ID NOs 3, 5, 19, 20, 22, 24, 28, and 29 as the bases for the design of the oligonucleotides. The oligonucleotides are selected to be approximately 10, 12, 15, 18, or more

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preferably 20 nucleotide residues in length, and to have a predicted hybridization temperature that is at least 37 degrees C. Preferably, the oligonucleotides are selected so that some will hybridize toward the 5' region of the mRNA molecule, others will hybridize to the coding region, and still others will hybridize to the 3' region of the mRNA molecule.

The oligonucleotides may be oligodeoxynucleotides, with phosphorothioate backbones (internucleoside linkages) throughout, or may have a variety of different types of internucleoside linkages. Generally, methods for the preparation, purification, and use of a variety of chemically modified oligonucleotides are described in U.S. Patent No. 5,948,680. Modified oligonucleosides may also be used in oligonucleotide synthesis, as well as mixed backbone compounds having, for instance, alternating MMI and P=O or P=S linkages, which are prepared as described in U.S. Pat. Nos. 5,378,825, 5,386,023, 5,489,677, 5,602,240 and 5,610,289. Formacetal- and thioformacetal-linked oligonucleosides may also be used and are prepared as described in U.S. Pat. Nos. 5,264,562 and 5,264,564; and ethylene oxide linked oligonucleosides may also be used and are prepared as described in U.S. Pat. No. 5,223,618. Peptide nucleic acids (PNAs) may be used as in the same manner as the oligonucleotides described above, and are prepared in accordance with any of the various procedures referred to in Peptide Nucleic Acids (PNA): Synthesis, Properties and Potential Applications, Bioorganic & Medicinal Chemistry, 1996, 4, 5-23; and U.S. Pat. Nos. 5,539,082, 5,700,922, and 5,719,262. Chimeric oligonucleotides, oligonucleosides, or mixed oligonucleotides/oligonucleosides of the invention are synthesized according to U.S. Pat. No. 5,623,065.

The effect of antisense compounds on target nucleic acid expression can be tested in any of a variety of cell types provided that the target nucleic acid is present at measurable levels. This can be routinely determined using, for example, PCR or Northern blot analysis. Preferably, the effect of several different oligonucleotides should be tested simultaneously, where the oligonucleotides hybridize to different portions of the target nucleic acid molecules, in order to identify the oligonucleotides producing the greatest degree of inhibition of expression of the target nucleic acid. Antisense modulation of HPR1 and/or HPR2 nucleic acid expression can be assayed in a variety of ways known in the art. For example, HPR1 and HPR2 mRNA levels can be quantitated by, e.g., Northern blot analysis, competitive polymerase chain reaction (PCR), or real-time PCR (RT-PCR). Real-time quantitative PCR is presently preferred. RNA analysis can be performed on total cellular RNA or poly(A)+ mRNA. Methods of RNA isolation and Northern blot analysis are taught in, for example, Ausubel, F. M. et al., Current Protocols in Molecular Biology, Volume 1, pp. 4.1.1-4.2.9 and 4.5.1-4.5.3, John Wiley & Sons, Inc., 1996. Real-time quantitative (PCR) can be conveniently accomplished using the commercially available ABI PRISM 7700 Sequence Detection System, available from PE-Applied Biosystems, Foster City, Calif. and used according to manufacturer's instructions. Other methods of quantitative PCR analysis are also known in the art. HPR1 and HPR2 protein levels can be quantitated in a variety of ways well known in the art, such as immunoprecipitation, Western blot analysis (immunoblotting), ELISA, or fluorescence-activated cell sorting (FACS). Antibodies directed to HPR1 and/or HPR2 polypeptides can be prepared via

conventional antibody generation methods such as those described herein. Immunoprecipitation methods, Western blot (immunoblot) analysis, and enzyme-linked immunosorbent assays (ELISA) are standard in the art (see, for example, Ausubel, F. M. et al., Current Protocols in Molecular Biology, Volume 2, pp. 10.16.1-10.16.11, 10.8.1-10.8.21, and 11.2.1-11.2.22, John Wiley & Sons, Inc., 1991).

All publications and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference. Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it will be readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

Sequences Presented in the Sequence Listing

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SEQ ID NO	Туре	Description
SEQ ID NO:1	Amino acid	Partial human HPR1 amino acid sequence
SEQ ID NO:2	Amino acid	Partial human HPR1 amino acid sequence
SEQ ID NO:3	Nucleotide	Human HPR1 cDNA sequence
SEQ ID NO:4	Amino acid	Human HPR1 amino acid sequence (745 amino acids)
SEQ ID NO:5	Nucleotide	Human HPR1 coding sequence
SEQ ID NO:6	Amino acid	Human LIF-R amino acid sequence (GenBank NP_002301)
SEQ ID NO:7	Amino acid	Human IL-12RB2 amino acid sequence (GenBank NP_001550)
SEQ ID NO:8	Amino acid	Human gp130 amino acid sequence (GenBank NP_002175)
SEQ ID NO:9	Amino acid	Human GCSFR amino acid sequence (SWISS-PROT Q99062)
SEQ ID NO:10	Amino acid	Portion of possible alternatively spliced form of human HPR1
SEQ ID NO:11	Amino acid	Portion of possible alternatively spliced form of human HPR1
SEQ ID NO:12	Amino acid	Mus musculus HPR1 amino acid sequence
SEQ ID NO:13	Amino acid	Possible 252-aa human HPR1 variant (WO 00/75314)
SEQ ID NO:14	Amino acid	Possible 652-aa human HPR1 variant (WO 00/75314)
SEQ ID NO:15	Amino acid	Possible 662-aa human HPR1 variant (WO 00/75314)
SEQ ID NO:16	Amino acid	Portion of possible alternatively spliced form of human HPR2
SEQ ID NO:17	Amino acid	Portion of possible alternatively spliced form of human HPR2
SEQ ID NO:18	Amino acid	Portion of possible alternatively spliced form of human HPR2
SEQ ID NO:19	Nucleotide	Human HPR2 cDNA sequence - exons 1 through 11
SEQ ID NO:20	Nucleotide	Human HPR2 coding sequence (encodes 629-aa form)
SEQ ID NO:21	Amino acid	Human HPR2 amino acid sequence (629 amino acids)
SEQ ID NO:22	Nucleotide	Human HPR2-ex8-ex9 coding sequence (encodes 565-aa form)
SEQ ID NO:23	Amino acid	Human HPR2-ex8-ex9 amino acid sequence (565 amino acids)
SEQ ID NO:24	Nucleotide	Human HPR2-ex9 coding sequence (encodes 356-aa form)
SEQ ID NO:25	Amino acid	Human HPR2-ex9 amino acid sequence (356 amino acids)
SEQ ID NO:26	Amino acid	Possible 384-aa human HPR2 variant (WO 00/73451)
SEQ ID NO:27	Amino acid	Mus musculus HPR2 amino acid sequence
SEQ ID NO:28	Nucleotide	Mus musculus HPR1 coding sequence
SEQ ID NO:29	Nucleotide	Mus musculus HPR2 coding sequence

5 What is claimed is:

**CLAIMS** 

1. An isolated polypeptide having HPR1 polypeptide activity comprising an amino acid sequence selected from the group consisting of:

(a) the amino acid sequence of SEQ ID NO:4;

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(b) an amino acid sequence selected from the group consisting of: amino acids 652 though 745 of SEQ ID NO:4, a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least 20 contiguous amino acids; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least 30 contiguous amino acids; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 that is at least 25% of the length of the sequence of amino acids 652 though 745 of SEQ ID NO:4; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 that is at least 50% of the length of the sequence of amino acids 652 though 745 of SEQ ID NO:4; and a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4; and a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4; and a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least eight contiguous amino acids and comprising at least one tyrosine residue;

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(c) an amino acid sequence comprising at least 8 amino acids and sharing amino acid identity with the amino acid sequences of (b), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 97.5%, at least 97.5%, at least 99.5%;

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an amino acid sequence comprising both an amino acid sequence of (b) or (c), and an amino acid sequence selected from the group consisting of: amino acids 1 through 55 of SEQ ID NO:1; amino acids 56 through 77 of SEO ID NO:1; amino acids 5 through 40 of SEO ID NO:2; amino acids 1 through 32 of SEQ ID NO:4; amino acids 1 through 241 of SEQ ID NO:4; amino acids 1 through 525 of SEQ ID NO:4; amino acids 20 through 32 of SEQ ID NO:4; amino acids 33 through 134 of SEQ ID NO:4; amino acids Xaa1 through Xaa2 of SEO ID NO:4, wherein Xaa1 is selected from the group consisting of amino acids 33 through 43 of SEQ ID NO:4 and Xaa2 is selected from the group consisting of amino acids 228 through 241 of SEQ ID NO:4; amino acids 33 through 238 of SEQ ID NO:4; amino acids 33 through 241 of SEQ ID NO:4; amino acids 33 through 525 of SEQ ID NO:4; amino acids 33 through 745 of SEQ ID NO:4; amino acids 44 through 94 of SEQ ID NO:4; amino acids 139 through 241 of SEQ ID NO:4; amino acids 242 through 326 of SEQ ID NO:4; amino acids 242 through 514 of SEO ID NO:4; amino acids 337 through 419 of SEQ ID NO:4; amino acids 433 through 514 of SEQ ID NO:4; amino acids 526 through 556 of SEQ ID NO:4; amino acids 533 through 552 of SEO ID NO:4; amino acids 553 through 745 of SEQ ID NO:4; amino acids 557 through 745 of SEQ ID NO:4; amino acids 563 through 573 of SEQ ID NO:4; amino acids 563 through 641 of SEQ ID NO:4; amino acids 567 through 581 of SEQ ID NO:4; amino acids 588 through 639 of SEQ ID NO:4; amino acids 631

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through 641 of SEQ ID NO:4; SEQ ID NO:10; and SEO ID NO:11:

5 (e) an amino acid sequence comprising both an amino acid sequence of (b) or (c), and a fragment of SEQ ID NO:4 comprising cytokine receptor domain amino acid sequences;

- (f) an allelic variant of any of (a)-(e); and
- (g) an amino acid sequence of (a)-(f), wherein a polypeptide comprising said amino acid sequence of (a)-(f) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (b)-(c).
  - 2. An isolated polypeptide having HPR1 polypeptide activity comprising an amino acid sequence selected from the group consisting of:
    - (a) the amino acid sequence of SEQ ID NO:12;

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- 15 (b) an amino acid sequence selected from the group consisting of: amino acids 633 though 726 of SEQ ID NO:12, a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 comprising at least 20 contiguous amino acids; a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 comprising at least 30 contiguous amino acids; a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 that is at least 25% of the length of the sequence of amino acids 633 though 726 of SEQ ID NO:12; a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 that is at least 50% of the length of the sequence of amino acids 633 though 726 of SEQ ID NO:12; and a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12; and a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 comprising at least eight contiguous amino acids and comprising at least one tyrosine residue;
  - (c) an amino acid sequence comprising at least 8 amino acids and sharing amino acid identity with the amino acid sequences of (b), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99.5%;
  - an amino acid sequence comprising both an amino acid sequence of (b) or (c), and an amino acid sequence selected from the group consisting of: amino acids 1 through 28 of SEQ ID NO:12; amino acids 1 through 224 of SEQ ID NO:12; amino acids 1 through 509 of SEQ ID NO:12; amino acids 13 through 28 of SEQ ID NO:12; amino acids 29 through 124 of SEQ ID NO:12; amino acids Xaa1 through Xaa2 of SEQ ID NO:12, wherein Xaa1 is selected from the group consisting of amino acids 29 through 39 of SEQ ID NO:12 and Xaa2 is selected from the group consisting of amino acids 211 through 224 of SEQ ID NO:12; amino acids 29 through 128 of SEQ ID NO:12; amino acids 29 through 509 of SEQ ID NO:12; amino acids 29 through 726 of SEQ ID NO:12; amino acids 129 through 224 of SEQ ID NO:12; amino acids 225 through 499 of SEQ ID NO:12; amino acids 320 through 403 of SEQ ID NO:12; amino acids 417 through 508 of SEQ ID NO:12; amino acids 510 through 533 of SEQ ID NO:12; amino acids 547 through 532 of SEQ ID NO:12; amino acids 534 through 726 of SEQ ID NO:12; amino acids 547 through 557 of

5 SEQ ID NO:12; amino acids 547 through 622 of SEQ ID NO:12; and amino acids 612 through 622 of SEQ ID NO:12;

- (e) an amino acid sequence comprising both an amino acid sequence of (b) or (c), and a fragment of SEQ ID NO:12 comprising cytokine receptor domain amino acid sequences;
  - (f) an allelic variant of any of (a)-(e); and
- 10 (g) an amino acid sequence of (a)-(f), wherein a polypeptide comprising said amino acid sequence of (a)-(f) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (b)-(c).
- 3. An isolated polypeptide having HPR2 polypeptide activity comprising an amino acid sequence selected from the group consisting of:
  - (a) SEQ ID NO:23;
  - (b) SEQ ID NO:25;

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- (c) an amino acid sequence selected from the group consisting of: an amino acid sequence comprising at least 20 contiguous amino acids of SEQ ID NO:23 and comprising the contiguous amino acids 318 and 319 of SEQ ID NO:23; and amino acids 349 through 356 of SEQ ID NO:25;
- (d) an amino acid sequence comprising at least 8 amino acids and sharing amino acid identity with the amino acid sequences of (c), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 97.5%, at least 97.5%, at least 99.5%;
- (e) an amino acid sequence comprising both an amino acid sequence of (c) or (d), and an amino acid sequence selected from the group consisting of: amino acids 1 through 177 of SEQ ID NO:16; amino acids 216 through 245 of SEQ ID NO:16; SEQ ID NO:17; and SEQ ID NO:18;
- (f) an amino acid sequence comprising both an amino acid sequence of (c) or (d), and an amino acid sequences of any of (a)-(b) comprising cytokine receptor domain amino acid sequences;
  - (g) an allelic variant of any of (a)-(f); and
- (h) an amino acid sequence of (a)-(g), wherein a polypeptide comprising said amino acid sequence of (a)-(g) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (c)-(d).
- 4. An isolated polypeptide having HPR2 polypeptide activity comprising an amino acid sequence selected from the group consisting of:
  - (a) SEQ ID NO:27;
- 40 (b) SEQ ID NO:27 from which amino acids 297 through 316 or amino acids 317 through 336 have been deleted;

5 (c) an amino acid sequence comprising 20 or more contiguous amino acids of (a) or (b); and

- (d) an amino acid sequence comprising 30 or more contiguous amino and sharing at least 90% amino acid identity with the amino acid sequences of (a)-(b).
- 10 5. An isolated nucleic acid encoding a polypeptide of any of claims 1 through 4.
  - 6. The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group consisting of:
    - (a) SEQ ID NO:3;
- 15 (b) SEQ ID NO:5;
  - (c) nucleotides 132 through 2366 of SEQ ID NO:3; and
  - (d) allelic variants of (a)-(c).
- 7. The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group consisting of nucleotides 1 through 137 of SEQ ID NO:3, nucleotides 138 through 228 of SEQ ID NO:3, nucleotides 229 through 346 of SEQ ID NO:3, nucleotides 347 through 528 of SEQ ID NO:3, nucleotides 529 through 680 of SEQ ID NO:3, nucleotides 681 through 846 of SEQ ID NO:3, nucleotides 847 through 926 of SEQ ID NO:3, nucleotides 927 through 1143 of SEQ ID NO:3, nucleotides 1144 through 1326 of SEQ ID NO:3, nucleotides 1327 through 1428 of SEQ ID NO:3, nucleotides 1429 through 1575 of SEQ ID NO:3, nucleotides 1576 through 1716 of SEQ ID NO:3, nucleotides 1717 through 1810 of SEQ ID NO:3, nucleotides 1811 through 1892 of SEQ ID NO:3, and nucleotides 1893 through 2480 of SEQ ID NO:3.
- 8. The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group consisting of:
  - (a) SEQ ID NO:28; and
  - (b) allelic variants of (a).
- 9. The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group consisting of:
  - (a) SEQ ID NO:19;
  - (b) SEQ ID NO:20;
  - (c) SEQ ID NO:22;
  - (d) SEQ ID NO:24; and
- 40 (d) allelic variants of (a)-(d).

5 10. The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group consisting of: nucleotides 107 through 175 of SEQ ID NO:19, nucleotides 107 through 478 of SEQ ID NO:19, nucleotides 107 through 1060 of SEQ ID NO:19, nucleotides 107 through 1099 of SEQ ID NO:19, nucleotides 107 through 1171 of SEQ ID NO:19, nucleotides 176 through 478 of SEQ ID NO:19, nucleotides 176 through 1099 of SEQ ID NO:19, nucleotides 176 through 1171 of SEQ ID NO:19, nucleotides 479 through 763 of SEQ ID NO:19, nucleotides 479 through 1099 of SEQ ID NO:19, nucleotides 503 through 1033 of SEQ ID NO:19, nucleotides 776 through 1066 of SEQ ID NO:19, nucleotides 776 through 1099 of SEQ ID NO:19, nucleotides 1232 through 1993 of SEQ ID NO:19, nucleotides 1283 through 1426 of SEQ ID NO:19, nucleotides 1283 through 1939 of SEQ ID NO:19, nucleotides 1295 through 1939

11. The nucleic acid of claim 5 comprising the nucleotide sequence of SEQ ID NO:29.

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ID NO:19.

- 12. An isolated genomic nucleic acid corresponding to the nucleic acid of any of claims 5 to 11.
  - 13. An expression vector comprising at least one nucleic acid according to any of claims 5 though 12.
- 14. A recombinant host cell comprising at least one nucleic acid according to any of claims 5 though 12.
  - 15. The recombinant host cell of claim 14, wherein the nucleic acid is integrated into the host cell genome.
- 30 16. A process for producing a polypeptide encoded by the nucleic acid of any of claims 5 though 12, comprising culturing a recombinant host cell under conditions promoting expression of said polypeptide, wherein the recombinant host cell comprises at least one nucleic acid according to any of claims 5 though 12.
- The process of claim 16 further comprising purifying said polypeptide.
  - 18. The polypeptide produced by the process of claim 17.
  - 19. An isolated antibody that binds to the polypeptide of any of claims 1 through 4 or claim 18.
  - 20. The antibody of claim 19 wherein the antibody is a monoclonal antibody.

- 5 21. The antibody of claim 19 wherein the antibody is a human antibody.
  - 22. The antibody of claim 19 wherein the antibody is a humanized antibody.
- 23. The antibody of claim 19 wherein the antibody inhibits the activity of the polypeptide of any of claims 1 through 4 or claim 18.
  - 24. A method of designing an inhibitor of the polypeptide of any of claims 1 through 4 or claim 18, the method comprising the steps of determining the three-dimensional structure of such polypeptide, analyzing the three-dimensional structure for the likely binding sites of substrates, synthesizing a molecule that incorporates a predicted reactive site, and determining the polypeptide-inhibiting activity of the molecule.
    - 25. A method for identifying compounds that alter HPR1 or HRP2 polypeptide activity comprising
- 20 (a) mixing a test compound with the polypeptide of any of claims 1 through 4 or claim 18; and
  - (b) determining whether the test compound alters the HPR1 or HRP2 polypeptide activity of said polypeptide.
- 25 26. A method for identifying compounds that inhibit the binding activity of HPR1 or HRP2 polypeptides comprising (a) mixing a test compound with the polypeptide of any of claims 1 through 4 or claim 18 and a binding partner of said polypeptide; and (b) determining whether the test compound inhibits the binding activity of said polypeptide.
- 30 27. A method for increasing ligand-binding activity comprising providing at least one compound selected from the group consisting of the polypeptide of any of claims 1 through 4 or claim 18 and agonists of said polypeptides.
- 28. The method of claim 27 wherein the method comprises increasing ligand-binding activity in a patient by administering to said patient at least one compound selected from the group consisting of the polypeptide of any of claims 1 through 4 or claim 18 and agonists of said polypeptides.
  - 29. A method for decreasing ligand-binding activity comprising providing at least one antagonist of the polypeptide of any of claims 1 through 4 or claim 18.

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5 30. The method of claim 29 wherein the method comprises decreasing ligand-binding activity in a patient by administering at least one antagonist of the polypeptide of any of claims 1 through 4 or claim 18 to said patient.

- 31. The method of claim 29 wherein the antagonist is an antibody that inhibits the activity of said polypeptide.
  - 32. A method for treating a cell proliferation condition comprising administering at least one compound selected from the group consisting of the polypeptide of any of claims 1 through 4 or claim 18 and agonists of said polypeptides.

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- 33. The method of claim 32 wherein the cell proliferation condition is selected from the group consisting of pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, and osteoporosis resulting from a lack of bone-forming cells.
- 20 34. A method for treating a cell proliferation condition comprising administering an antagonist of the polypeptide of any of claims 1 through 4 or claim 18.
- 35. The method of claim 34 wherein the cell proliferation condition is selected from the group consisting of leukemia, tumour metastasis, and osteoporosis resulting from an excess of bone-resorbing cells.

## SEQUENCE LISTING

<110> Immunex Corporation Cosman, David J. Mosley, Bruce A. Bird, Timothy A. DuBose, Robert F. Wiley, Steven R.

<120> HEMATOPOIETIN RECEPTORS HPR1 AND HPR2

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435

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Leu 465	Cys	Glu	Ile	Glu	Ile 470	Lys	Lys	Ser	Asn	Ser 475	Val	Gln	Glu	Gln	Arg 480
Asn	Val	Thr	Ile	Ьув 485	G1y	Val	G1u	Asn	Ser 490	Ser	Tyr	Leu	Val	Ala 495	Leu
Asp	Lys	Leu	Asn 500	Pro	Tyr	Thr	Leu	Tyr 505	Thr	Phe	Arg	Ile	Arg 510	Cys	Ser
Thr	Glu	Thr 515	Phe	Trp	ГЛЗ	Trp	Ser 520	Lys	Trp	Ser	Asn	Lys 525	Lys	Gln	His
Leu	Thr 530	Thr	Glu	Ala	Ser	Pro 535	Ser	Lys	Gly	Pro	Asp 540	Thr	Trp	Arg	Glu
Trp 545	Ser	Ser	Asp	Gly	Lys 550	Asn	Leu	Ile	Ile	Tyr 555	Trp	Lys	Pro	Leu	Pro 560
Ile	Asn	Glu	Ala	Asn 565	Gly	Lys	Ile	Leu	Ser 570	Tyr	Asn	Val	Ser	Cys 575	Ser
Ser	Asp	Glu	Glu 580	Thr	Gln	Ser	Leu	Ser 585	Glu	Ile	Pro	Asp	Pro 590	Gln	His
Lys	Ala	Glu 595	Ile	Arg	Leu	Asp	Lys 600	Asn	Asp	Tyr	Ile	Ile 605	Ser	Val	Val
Ala	Lys 610	Asn	Ser	Val	Gly	Ser 615	Ser	Pro	Pro	Ser	Lys 620	Ile	Ala	Ser	Met
Glu 625	Ile	Pro	Asn	Asp	Asp 630	Leu	Lys	Ile	Glu	Gln 635	Val	Val	Gly	Met	Gly 640
Lys	Gly	Ile	Leu	Leu 645	Thr	Trp	His	Tyr	Asp 650	Pro	Asn	Met	Thr	Cys 655	Asp
Tyr	Val	Ile	Lys 660	Trp	Cys	Asn	Ser	Ser 665	Arg	Ser	Glu	Pro	Cys 670	Leu	Met
Asp	Trp	Arg 675	Lys	Val	Pro	Ser	Asn 680	Ser	Thr	Glu	Thr	Val 685	Ile	Glu	Ser

Asp Glu Phe Arg Pro Gly Ile Arg Tyr Asn Phe Phe Leu Tyr Gly Cys Arg Asn Gln Gly Tyr Gln Leu Leu Arg Ser Met Ile Gly Tyr Ile Glu 710 715 705 Glu Leu Ala Pro Ile Val Ala Pro Asn Phe Thr Val Glu Asp Thr Ser Ala Asp Ser Ile Leu Val Lys Trp Glu Asp Ile Pro Val Glu Leu 745 Arg Gly Phe Leu Arg Gly Tyr Leu Phe Tyr Phe Gly Lys Gly Glu Arg 755 760 Asp Thr Ser Lys Met Arg Val Leu Glu Ser Gly Arg Ser Asp Ile Lys 775 Val Lys Asn Ile Thr Asp Ile Ser Gln Lys Thr Leu Arg Ile Ala Asp Leu Gln Gly Lys Thr Ser Tyr His Leu Val Leu Arg Ala Tyr Thr Asp 805 810 Gly Val Gly Pro Glu Lys Ser Met Tyr Val Val Thr Lys Glu Asn 825 820 Ser Val Gly Leu Ile Ile Ala Ile Leu Ile Pro Val Ala Val Ala Val 835 840 Ile Val Gly Val Val Thr Ser Ile Leu Cys Tyr Arg Lys Arg Glu Trp Ile Lys Glu Thr Phe Tyr Pro Asp Ile Pro Asn Pro Glu Asn Cys Lys 870 875 Ala Leu Gln Phe Gln Lys Ser Val Cys Glu Gly Ser Ser Ala Leu Lys Thr Leu Glu Met Asn Pro Cys Thr Pro Asn Asn Val Glu Val Leu Glu 905 Thr Arg Ser Ala Phe Pro Lys Ile Glu Asp Thr Glu Ile Ile Ser Pro 915 920 925

Val Ala Glu Arg Pro Glu Asp Arg Ser Asp Ala Glu Pro Glu Asn His 930 935 940

Val Val Val Ser Tyr Cys Pro Pro Ile Ile Glu Glu Glu Ile Pro Asn 945 950 955 960

Pro Ala Ala Asp Glu Ala Gly Gly Thr Ala Gln Val Ile Tyr Ile Asp 965 970 975

Val Gln Ser Met Tyr Gln Pro Gln Ala Lys Pro Glu Glu Glu Glu Glu 980 985 990

Asn Asp Pro Val Gly Gly Ala Gly Tyr Lys Pro Gln Met His Leu Pro 995 1000 1005

Ile Asn Ser Thr Val Glu Asp Ile Ala Ala Glu Glu Asp Leu Asp 1010 1015 1020

Lys Thr Ala Gly Tyr Arg Pro Gln Ala Asn Val Asn Thr Trp Asn 1025 1030 1035

Leu Val Ser Pro Asp Ser Pro Arg Ser Ile Asp Ser Asn Ser Glu 1040 1045 1050

Ile Val Ser Phe Gly Ser Pro Cys Ser Ile Asn Ser Arg Gln Phe 1055 1060 1065

Leu Ile Pro Pro Lys Asp Glu Asp Ser Pro Lys Ser Asn Gly Gly 1070 1075 1080

<210> 7

<211> 979

<212> PRT

<213> Homo sapiens

<400> 7

Met Ala Leu Phe Ala Val Phe Gln Thr Thr Phe Phe Leu Thr Leu Leu 1 5 10 15

Ser Leu Arg Thr Tyr Gln Ser Glu Val Leu Ala Glu Arg Leu Pro Leu 20 25 30

Thr Pro Val Ser Leu Lys Val Ser Thr Asn Ser Thr Arg Gln Ser Leu 35 40 45

His Leu Gln Trp Thr Val His Asn Leu Pro Tyr His Gln Glu Leu Lys Met Val Phe Gln Ile Gln Ile Ser Arg Ile Glu Thr Ser Asn Val Ile 70 65 75 Trp Val Gly Asn Tyr Ser Thr Thr Val Lys Trp Asn Gln Val Leu His 85 Trp Ser Trp Glu Ser Glu Leu Pro Leu Glu Cys Ala Thr His Phe Val 105 Arg Ile Lys Ser Leu Val Asp Asp Ala Lys Phe Pro Glu Pro Asn Phe 115 120 Trp Ser Asn Trp Ser Ser Trp Glu Glu Val Ser Val Gln Asp Ser Thr 135 Gly Gln Asp Ile Leu Phe Val Phe Pro Lys Asp Lys Leu Val Glu Glu 155 160 150 Gly Thr Asn Val Thr Ile Cys Tyr Val Ser Arg Asn Ile Gln Asn Asn 165 170 Val Ser Cys Tyr Leu Glu Gly Lys Gln Ile His Gly Glu Gln Leu Asp 180 185 Pro His Val Thr Ala Phe Asn Leu Asn Ser Val Pro Phe Ile Arg Asn 195 200 205 Lys Gly Thr Asn Ile Tyr Cys Glu Ala Ser Gln Gly Asn Val Ser Glu 210 215 Gly Met Lys Gly Ile Val Leu Phe Val Ser Lys Val Leu Glu Glu Pro 225 230 235 240 Lys Asp Phe Ser Cys Glu Thr Glu Asp Phe Lys Thr Leu His Cys Thr 245 Trp Asp Pro Gly Thr Asp Thr Ala Leu Gly Trp Ser Lys Gln Pro Ser 260 265 Gln Ser Tyr Thr Leu Phe Glu Ser Phe Ser Gly Glu Lys Lys Leu Cys 275 280 285

Thr His Lys Asn Trp Cys Asn Trp Gln Ile Thr Gln Asp Ser Gln Glu

Thr Tyr Asn Phe Thr Leu Ile Ala Glu Asn Tyr Leu Arg Lys Arg Ser Val Asn Ile Leu Phe Asn Leu Thr His Arg Val Tyr Leu Met Asn Pro Phe Ser Val Asn Phe Glu Asn Val Asn Ala Thr Asn Ala Ile Met Thr Trp Lys Val His Ser Ile Arg Asn Asn Phe Thr Tyr Leu Cys Gln Ile Glu Leu His Gly Glu Gly Lys Met Met Gln Tyr Asn Val Ser Ile Lys Val Asn Gly Glu Tyr Phe Leu Ser Glu Leu Glu Pro Ala Thr Glu Tyr Met Ala Arg Val Arg Cys Ala Asp Ala Ser His Phe Trp Lys Trp Ser Glu Trp Ser Gly Gln Asn Phe Thr Thr Leu Glu Ala Ala Pro Ser Glu Ala Pro Asp Val Trp Arg Ile Val Ser Leu Glu Pro Gly Asn His Thr Val Thr Leu Phe Trp Lys Pro Leu Ser Lys Leu His Ala Asn Gly Lys Ile Leu Phe Tyr Asn Val Val Val Glu Asn Leu Asp Lys Pro Ser Ser Ser Glu Leu His Ser Ile Pro Ala Pro Ala Asn Ser Thr Lys Leu Ile Leu Asp Arg Cys Ser Tyr Gln Ile Cys Val Ile Ala Asn Asn Ser Val Gly Ala Ser Pro Ala Ser Val Ile Val Ile Ser Ala Asp Pro Glu Asn 

Lys Glu Val Glu Glu Glu Arg Ile Ala Gly Thr Glu Gly Gly Phe Ser 535 Leu Ser Trp Lys Pro Gln Pro Gly Asp Val Ile Gly Tyr Val Val Asp 545 550 555 Trp Cys Asp His Thr Gln Asp Val Leu Gly Asp Phe Gln Trp Lys Asn 570 Val Gly Pro Asn Thr Thr Ser Thr Val Ile Ser Thr Asp Ala Phe Arg 580 585 Pro Gly Val Arg Tyr Asp Phe Arg Ile Tyr Gly Leu Ser Thr Lys Arg Ile Ala Cys Leu Leu Glu Lys Lys Thr Gly Tyr Ser Gln Glu Leu Ala 615 620 Pro Ser Asp Asn Pro His Val Leu Val Asp Thr Leu Thr Ser His Ser 630 635 Phe Thr Leu Ser Trp Lys Asp Tyr Ser Thr Glu Ser Gln Pro Gly Phe Ile Gln Gly Tyr His Val Tyr Leu Lys Ser Lys Ala Arg Gln Cys His 660 665 670 Pro Arg Phe Glu Lys Ala Val Leu Ser Asp Gly Ser Glu Cys Cys Lys 675 680 Tyr Lys Ile Asp Asn Pro Glu Glu Lys Ala Leu Ile Val Asp Asn Leu 690 695 Lys Pro Glu Ser Phe Tyr Glu Phe Phe Ile Thr Pro Phe Thr Ser Ala 705 710 715 720 Gly Glu Gly Pro Ser Ala Thr Phe Thr Lys Val Thr Thr Pro Asp Glu 725 730 His Ser Ser Met Leu Ile His Ile Leu Leu Pro Met Val Phe Cys Val 740 745 Leu Leu Ile Met Val Met Cys Tyr Leu Lys Ser Gln Trp Ile Lys Glu 755 760 765 Thr Cys Tyr Pro Asp Ile Pro Asp Pro Tyr Lys Ser Ser Ile Leu Ser

775 780 770

Leu Ile Lys Phe Lys Glu Asn Pro His Leu Ile Ile Met Asn Val Ser 790 795

Asp Cys Ile Pro Asp Ala Ile Glu Val Val Ser Lys Pro Glu Gly Thr

Lys Ile Gln Phe Leu Gly Thr Arg Lys Ser Leu Thr Glu Thr Glu Leu 825

Thr Lys Pro Asn Tyr Leu Tyr Leu Pro Thr Glu Lys Asn His Ser 835 840

Gly Pro Gly Pro Cys Ile Cys Phe Glu Asn Leu Thr Tyr Asn Gln Ala 855 850

Ala Ser Asp Ser Gly Ser Cys Gly His Val Pro Val Ser Pro Lys Ala 875

Pro Ser Met Leu Gly Leu Met Thr Ser Pro Glu Asn Val Leu Lys Ala

Leu Glu Lys Asn Tyr Met Asn Ser Leu Gly Glu Ile Pro Ala Gly Glu 900 905 910

Thr Ser Leu Asn Tyr Val Ser Gln Leu Ala Ser Pro Met Phe Gly Asp

Lys Asp Ser Leu Pro Thr Asn Pro Val Glu Ala Pro His Cys Ser Glu 935

Tyr Lys Met Gln Met Ala Val Ser Leu Arg Leu Ala Leu Pro Pro 950 955

Thr Glu Asn Ser Ser Leu Ser Ser Ile Thr Leu Leu Asp Pro Gly Glu 970

His Tyr Cys

<210> 8

<211> 918 <212> PRT

<213> Homo sapiens

<400> 8

Met Leu Thr Leu Gln Thr Trp Val Val Gln Ala Leu Phe Ile Phe Leu Thr Thr Glu Ser Thr Gly Glu Leu Leu Asp Pro Cys Gly Tyr Ile Ser 20 25 30 Pro Glu Ser Pro Val Val Gln Leu His Ser Asn Phe Thr Ala Val Cys Val Leu Lys Glu Lys Cys Met Asp Tyr Phe His Val Asn Ala Asn Tyr 50 Ile Val Trp Lys Thr Asn His Phe Thr Ile Pro Lys Glu Gln Tyr Thr 70 Ile Ile Asn Arg Thr Ala Ser Ser Val Thr Phe Thr Asp Ile Ala Ser 85 Leu Asn Ile Gln Leu Thr Cys Asn Ile Leu Thr Phe Gly Gln Leu Glu 105 Gln Asn Val Tyr Gly Ile Thr Ile Ile Ser Gly Leu Pro Pro Glu Lys Pro Lys Asn Leu Ser Cys Ile Val Asn Glu Gly Lys Lys Met Arg Cys 130 135 140 Glu Trp Asp Gly Gly Arg Glu Thr His Leu Glu Thr Asn Phe Thr Leu 150 Lys Ser Glu Trp Ala Thr His Lys Phe Ala Asp Cys Lys Ala Lys Arg Asp Thr Pro Thr Ser Cys Thr Val Asp Tyr Ser Thr Val Tyr Phe Val 185 Asn Ile Glu Val Trp Val Glu Ala Glu Asn Ala Leu Gly Lys Val Thr Ser Asp His Ile Asn Phe Asp Pro Val Tyr Lys Val Lys Pro Asn Pro 215 Pro His Asn Leu Ser Val Ile Asn Ser Glu Glu Leu Ser Ser Ile Leu 225 230 235

Lys Leu Thr Trp Thr Asn Pro Ser Ile Lys Ser Val Ile Ile Leu Lys Tyr Asn Ile Gln Tyr Arg Thr Lys Asp Ala Ser Thr Trp Ser Gln Ile Pro Pro Glu Asp Thr Ala Ser Thr Arg Ser Ser Phe Thr Val Gln Asp Leu Lys Pro Phe Thr Glu Tyr Val Phe Arg Ile Arg Cys Met Lys Glu Asp Gly Lys Gly Tyr Trp Ser Asp Trp Ser Glu Glu Ala Ser Gly Ile Thr Tyr Glu Asp Arg Pro Ser Lys Ala Pro Ser Phe Trp Tyr Lys Ile Asp Pro Ser His Thr Gln Gly Tyr Arg Thr Val Gln Leu Val Trp Lys Thr Leu Pro Pro Phe Glu Ala Asn Gly Lys Ile Leu Asp Tyr Glu Val Thr Leu Thr Arg Trp Lys Ser His Leu Gln Asn Tyr Thr Val Asn Ala Thr Lys Leu Thr Val Asn Leu Thr Asn Asp Arg Tyr Leu Ala Thr Leu Thr Val Arg Asn Leu Val Gly Lys Ser Asp Ala Ala Val Leu Thr Ile Pro Ala Cys Asp Phe Gln Ala Thr His Pro Val Met Asp Leu Lys Ala Phe Pro Lys Asp Asn Met Leu Trp Val Glu Trp Thr Thr Pro Arg Glu Ser Val Lys Lys Tyr Ile Leu Glu Trp Cys Val Leu Ser Asp Lys Ala Pro Cys Ile Thr Asp Trp Gln Glu Asp Gly Thr Val His Arg Thr Tyr Leu Arg Gly Asn Leu Ala Glu Ser Lys Cys Tyr Leu Ile Thr Val

				485					490					495	
Thr	Pro	Val	Tyr 500	Ala	Asp	Gly	Pro	Gly 505	Ser	Pro	Glu	Ser	Ile 510	Lys	Ala
Tyr	Leu	Lys 515	Gln	Ala	Pro	Pro	Ser 520	Lys	Gly	Pro	Thr	Val 525	Arg	Thr	Lys
Lys	Val 530	Gly	Lys	Asn	Glu	Ala 535	Val	Leu	Glu	Trp	Asp 540	Gln	Leu	Pro	Val
Asp 545	Val	Gln	Asn	Gly	Phe 550	Ile	Arg	Asn	Tyr	Thr 555	Ile	Phe	Tyr	Arg	Thr 560
Ile	Ile	Gly	Asn	Glu 565	Thr	Ala	Val	Asn	Val 570	Asp	Ser	Ser	His	Thr 575	Glu
Tyr	Thr	Leu	Ser 580	Ser	Leu	Thr	Ser	Asp 585	Thr	Leu	Туг	Met	Val 590	Arg	Met
Ala	Ala	Tyr 595	Thr	qaA	Glu	Gly	Gly 600	Lys	Asp	Gly	Pro	Glu 605	Phe	Thr	Phe
Thr	Thr 610	Pro	Ľys	Phe	Ala	Gln 615	Gly	Glu	Ile	Glu	Ala 620	Ile	Val	Val	Pro
Val 625	Cys	Leu	Ala	Phe	Leu 630	Leu	Thr	Thr	Leu	Leu 635	Gly	Val	Leu	Phe	Cys 640
Phe	Asn	Lys	Arg	Asp 645	Leu	Ile	Lys	Lys	His 650	Ile	Trp	Pro	Asn	Val 655	Pro
Asp	Pro	Ser	Ъуз 660	Ser	His	Ile	Ala	Gln 665	Trp	Ser	Pro	His	Thr 670	Pro	Pro
Arg	His	Asn 675	Phe	Asn	Ser	Lys	Asp 680	Gln	Met	Tyr	Ser	Asp 685	Gly	Asn	Phe
Thr	Asp 690	Val	Ser	Val	Val	G1u 695	Ile	Glu	Ala	Asn	Asp 700	Lys	Lys	Pro	Phe
Pro 705	Glu	Asp	Leu	Lys	Ser 710	Leu	Asp	Leu	Phe	Lys 715	Lys	Glu	Lys	Ile	Asn 720
Thr	Glu	Gly	His	Ser 725	Ser	Gly	Ile	Gly	Gly 730	Ser	Ser	Cys	Met	Ser 735	Ser

Ser Arg Pro Ser Ile Ser Ser Ser Asp Glu Asn Glu Ser Ser Gln Asn 740 745 750

Thr Ser Ser Thr Val Gln Tyr Ser Thr Val Val His Ser Gly Tyr Arg
755 760 765

His Gln Val Pro Ser Val Gln Val Phe Ser Arg Ser Glu Ser Thr Gln 770 780

Pro Leu Leu Asp Ser Glu Glu Arg Pro Glu Asp Leu Gln Leu Val Asp 785 790 795 800

His Val Asp Gly Gly Asp Gly Ile Leu Pro Arg Gln Gln Tyr Phe Lys 805 810 815

Gln Asn Cys Ser Gln His Glu Ser Ser Pro Asp Ile Ser His Phe Glu 820 825 830

Arg Ser Lys Gl<br/>n Val Ser Ser Val As<br/>n Glu Glu Asp Phe Val Arg Leu 835  $840 \,$  845

Lys Gln Gln Ile Ser Asp His Ile Ser Gln Ser Cys Gly Ser Gly Gln 850 855 860

Met Lys Met Phe Gln Glu Val Ser Ala Ala Asp Ala Phe Gly Pro Gly 865 870 875 880

Thr Glu Gly Gln Val Glu Arg Phe Glu Thr Val Gly Met Glu Ala Ala 885 890 895

Thr Asp Glu Gly Met Pro Lys Ser Tyr Leu Pro Gln Thr Val Arg Gln 900 905 910

Gly Gly Tyr Met Pro Gln 915

<210> 9

<211> 836

<212> PRT

<213> Homo sapiens

<400> 9

Met Ala Arg Leu Gly Asn Cys Ser Leu Thr Trp Ala Ala Leu Ile Ile 1  $\phantom{\bigg|}$  5  $\phantom{\bigg|}$  10  $\phantom{\bigg|}$  15

Leu Leu Pro Gly Ser Leu Glu Glu Cys Gly His Ile Ser Val Ser Ala Pro Ile Val His Leu Gly Asp Pro Ile Thr Ala Ser Cys Ile Ile 40 Lys Gln Asn Cys Ser His Leu Asp Pro Glu Pro Gln Ile Leu Trp Arg 55 Leu Gly Ala Glu Leu Gln Pro Gly Gly Arg Gln Gln Arg Leu Ser Asp Gly Thr Gln Glu Ser Ile Ile Thr Leu Pro His Leu Asn His Thr Gln 90 Ala Phe Leu Ser Cys Cys Leu Asn Trp Gly Asn Ser Leu Gln Ile Leu Asp Gln Val Glu Leu Arg Ala Gly Tyr Pro Pro Ala Ile Pro His Asn 120 Leu Ser Cys Leu Met Asn Leu Thr Thr Ser Ser Leu Ile Cys Gln Trp 135 Glu Pro Gly Pro Glu Thr His Leu Pro Thr Ser Phe Thr Leu Lys Ser Phe Lys Ser Arg Gly Asn Cys Gln Thr Gln Gly Asp Ser Ile Leu Asp 170 165 Cys Val Pro Lys Asp Gly Gln Ser His Cys Cys Ile Pro Arg Lys His , 180 Leu Leu Tyr Gln Asn Met Gly Ile Trp Val Gln Ala Glu Asn Ala 195 200 Leu Gly Thr Ser Met Ser Pro Gln Leu Cys Leu Asp Pro Met Asp Val 210 215 Val Lys Leu Glu Pro Pro Met Leu Arg Thr Met Asp Pro Ser Pro Glu 225 235 240 Ala Ala Pro Pro Gln Ala Gly Cys Leu Gln Leu Cys Trp Glu Pro Trp Gln Pro Gly Leu His Ile Asn Gln Lys Cys Glu Leu Arg His Lys Pro

260 265 270 Gln Arg Gly Glu Ala Ser Trp Ala Leu Val Gly Pro Leu Pro Leu Glu 275 280 Ala Leu Gln Tyr Glu Leu Cys Gly Leu Leu Pro Ala Thr Ala Tyr Thr 295 Leu Gln Ile Arg Cys Ile Arg Trp Pro Leu Pro Gly His Trp Ser Asp 305 310 315 Trp Ser Pro Ser Leu Glu Leu Arg Thr Thr Glu Arg Ala Pro Thr Val 325 330 Arg Leu Asp Thr Trp Trp Arg Gln Arg Gln Leu Asp Pro Arg Thr Val 340 345 Gln Leu Phe Trp Lys Pro Val Pro Leu Glu Glu Asp Ser Gly Arg Ile Gln Gly Tyr Val Val Ser Trp Arg Pro Ser Gly Gln Ala Gly Ala Ile Leu Pro Leu Cys Asn Thr Thr Glu Leu Ser Cys Thr Phe His Leu Pro 385 390 395 Ser Glu Ala Gln Glu Val Ala Leu Val Ala Tyr Asn Ser Ala Gly Thr 410 Ser Arg Pro Thr Pro Val Val Phe Ser Glu Ser Arg Gly Pro Ala Leu 425 Thr Arg Leu His Ala Met Ala Arg Asp Pro His Ser Leu Trp Val Gly 435 440 Trp Glu Pro Pro Asn Pro Trp Pro Gln Gly Tyr Val Ile Glu Trp Gly 455 460 Leu Gly Pro Pro Ser Ala Ser Asn Ser Asn Lys Thr Trp Arg Met Glu 470 475 Gln Asn Gly Arg Ala Thr Gly Phe Leu Leu Lys Glu Asn Ile Arg Pro 485 490 Phe Gln Leu Tyr Glu Ile Ile Val Thr Pro Leu Tyr Gln Asp Thr Met 500 505 510

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Gly Pro Ser Gln His Val Tyr Ala Tyr Ser Gln Glu Met Ala Pro Ser 520 His Ala Pro Glu Leu His Leu Lys His Ile Gly Lys Thr Trp Ala Gln 535 Leu Glu Trp Val Pro Glu Pro Pro Glu Leu Gly Lys Ser Pro Leu Thr 545 550 555 His Tyr Thr Ile Phe Trp Thr Asn Ala Gln Asn Gln Ser Phe Ser Ala 565 570 Ile Leu Asn Ala Ser Ser Arg Gly Phe Val Leu His Gly Leu Glu Pro Ala Ser Leu Tyr His Ile His Leu Met Ala Ala Ser Gln Ala Gly Ala 600 Thr Asn Ser Thr Val Leu Thr Leu Met Thr Leu Thr Pro Glu Gly Ser Glu Leu His Ile Ile Leu Gly Leu Phe Gly Leu Leu Leu Leu Thr 635 Cys Leu Cys Gly Thr Ala Trp Leu Cys Cys Ser Pro Asn Arg Lys Asn 650 645 Pro Leu Trp Pro Ser Val Pro Asp Pro Ala His Ser Ser Leu Gly Ser 660 665 670 Trp Val Pro Thr Ile Met Glu Glu Asp Ala Phe Gln Leu Pro Gly Leu 675 680 Gly Thr Pro Pro Ile Thr Lys Leu Thr Val Leu Glu Glu Asp Glu Lys 690 695 Lys Pro Val Pro Trp Glu Ser His Asn Ser Ser Glu Thr Cys Gly Leu 705 710 715 720 Pro Thr Leu Val Gln Thr Tyr Val Leu Gln Gly Asp Pro Arg Ala Val Ser Thr Gln Pro Gln Ser Gln Ser Gly Thr Ser Asp Gln Val Leu Tyr 740 745

Gly Gln Leu Leu Gly Ser Pro Thr Ser Pro Gly Pro Gly His Tyr Leu Arg Cys Asp Ser Thr Gln Pro Leu Leu Ala Gly Leu Thr Pro Ser Pro 775 770 780 Lys Ser Tyr Glu Asn Leu Trp Phe Gln Ala Ser Pro Leu Gly Thr Leu 790 795 Val Thr Pro Ala Pro Ser Gln Glu Asp Asp Cys Val Phe Gly Pro Leu 810 Leu Asn Phe Pro Leu Leu Gln Gly Ile Arg Val His Gly Met Glu Ala 820 825 Leu Gly Ser Phe 835 <210> 10 <211> 7 <212> PRT <213> Homo sapiens <400> 10 Trp Lys Ser Thr Ser Val Lys 1. 5 <210> 11 <211> 15 <212> PRT <213> Homo sapiens <400> 11 Glu Gly Lys Leu Leu Pro Ala Ile Pro Val Leu Ser Ala Leu Lys 10 <210> 12 <211> 726 <212> PRT <213> Mus musculus <400> 12 Met Lys Pro Leu Gly Val Asn Ala Gly Ile Met Trp Thr Leu Ala Leu Trp Ala Phe Ser Phe Leu Cys Lys Phe Ser Leu Ala Val Leu Pro Thr 2.0 25 3.0

Lys Pro Glu Asn Ile Ser Cys Val Phe Tyr Phe Asp Arg Asn Leu Thr 40 Cys Thr Trp Arg Pro Glu Lys Glu Thr Asn Asp Thr Ser Tyr Ile Val 50 55 Thr Leu Thr Tyr Ser Tyr Gly Lys Ser Asn Tyr Ser Asp Asn Ala Thr 70 75 Glu Ala Ser Tyr Ser Phe Pro Arg Ser Cys Ala Met Pro Pro Asp Ile 90 95 Cys Ser Val Glu Val Gln Ala Gln Asn Gly Asp Gly Lys Val Lys Ser 100 105 Asp Ile Thr Tyr Trp His Leu Ile Ser Ile Ala Lys Thr Glu Pro Pro 115 120 Ile Ile Leu Ser Val Asn Pro Ile Cys Asn Arg Met Phe Gln Ile Gln 130 135 Trp Lys Pro Arg Glu Lys Thr Arg Gly Phe Pro Leu Val Cys Met Leu Arg Phe Arg Thr Val Asn Ser Ser Arg Trp Thr Glu Val Asn Phe Glu 165 170 Asn Cys Lys Gln Val Cys Asn Leu Thr Gly Leu Gln Ala Phe Thr Glu 180 185 Tyr Val Leu Ala Leu Arg Phe Arg Phe Asn Asp Ser Arg Tyr Trp Ser 200 Lys Trp Ser Lys Glu Glu Thr Arg Val Thr Met Glu Glu Val Pro His 215 Val Leu Asp Leu Trp Arg Ile Leu Glu Pro Ala Asp Met Asn Gly Asp 230 235 Arg Lys Val Arg Leu Leu Trp Lys Lys Ala Arg Gly Ala Pro Val Leu Glu Lys Thr Phe Gly Tyr His Ile Gln Tyr Phe Ala Glu Asn Ser Thr 265

Asn Leu Thr Glu Ile Asn Asn Ile Thr Thr Gln Gln Tyr Glu Leu Leu Leu Met Ser Gln Ala His Ser Val Ser Val Thr Ser Phe Asn Ser Leu Gly Lys Ser Gln Glu Thr Ile Leu Arg Ile Pro Asp Val His Glu Lys Thr Phe Gln Tyr Ile Lys Ser Met Gln Ala Tyr Ile Ala Glu Pro Leu Leu Val Val Asn Trp Gln Ser Ser Ile Pro Ala Val Asp Thr Trp Ile Val Glu Trp Leu Pro Glu Ala Ala Met Ser Lys Phe Pro Ala Leu Ser Trp Glu Ser Val Ser Gln Val Thr Asn Trp Thr Ile Glu Gln Asp Lys Leu Lys Pro Phe Thr Cys Tyr Asn Ile Ser Val Tyr Pro Val Leu Gly His Arg Val Gly Glu Pro Tyr Ser Ile Gln Ala Tyr Ala Lys Glu Gly Thr Pro Leu Lys Gly Pro Glu Thr Arg Val Glu Asn Ile Gly Leu Arg Thr Ala Thr Ile Thr Trp Lys Glu Ile Pro Lys Ser Ala Arg Asn Gly Phe Ile Asn Asn Tyr Thr Val Phe Tyr Gln Ala Glu Gly Gly Lys Glu Leu Ser Lys Thr Val Asn Ser His Ala Leu Gln Cys Asp Leu Glu Ser Leu Thr Arg Arg Thr Ser Tyr Thr Val Trp Val Met Ala Ser Thr Arg Ala Gly Gly Thr Asn Gly Val Arg Ile Asn Phe Lys Thr Leu Ser Ile Ser Val Phe Glu Val Val Leu Leu Thr Ser Leu Val Gly Gly Leu

520 525 515

Leu Leu Ser Ile Lys Thr Val Thr Phe Gly Leu Arg Lys Pro Asn 530 535 540

Arg Leu Thr Pro Leu Cys Cys Pro Asp Val Pro Asn Pro Ala Glu Ser 545 550 555

Ser Leu Ala Thr Trp Leu Gly Asp Gly Phe Lys Lys Ser Asn Met Lys 565 570

Glu Thr Gly Asn Ser Gly Asn Thr Glu Asp Val Val Leu Lys Pro Cys

Pro Val Pro Ala Asp Leu Ile Asp Lys Leu Val Val Asn Phe Glu Asn 595 600 605

Phe Leu Glu Val Val Leu Thr Glu Glu Ala Gly Lys Gly Gln Ala Ser

Ile Leu Gly Gly Glu Ala Asn Glu Tyr Val Thr Ser Pro Ser Arg Pro 625

Asp Gly Pro Pro Gly Lys Ser Phe Lys Glu Pro Ser Ile Leu Thr Glu 645 650

Val Ala Ser Glu Asp Ser His Ser Thr Cys Ser Arg Met Ala Asp Glu 660 665

Ala Tyr Ser Glu Leu Ala Arg Gln Pro Ser Ser Ser Cys Gln Ser Pro 680

Gly Leu Ser Pro Pro Arg Glu Asp Gln Ala Gln Asn Pro Tyr Leu Lys 695

Asn Ser Val Thr Thr Arg Glu Phe Leu Val His Glu Asn Ile Pro Glu 715 710

His Ser Lys Gly Glu Val 725

<210> 13 <211> 252

<212> PRT

<213> Homo sapiens

<400> 13

Met Lys Leu Ser Pro Gln Pro Ser Cys Val Asn Leu Gly Met Met Trp Thr Trp Ala Leu Trp Met Leu Pro Ser Leu Cys Lys Phe Ser Leu Ala 25 20 Ala Leu Pro Ala Lys Pro Glu Asn Ile Ser Cys Val Tyr Tyr Tyr Arg Lys Asn Leu Thr Cys Thr Trp Ser Pro Gly Lys Glu Thr Ser Tyr Thr Gln Tyr Thr Val Lys Arg Thr Tyr Ala Phe Gly Glu Lys His Asp Asn 70 Cys Thr Thr Asn Ser Ser Thr Ser Glu Asn Arg Ala Ser Cys Ser Phe Phe Leu Pro Arg Ile Thr Ile Pro Asp Asn Tyr Thr Ile Glu Val Glu 105 Ala Glu Asn Gly Asp Gly Val Ile Lys Ser His Met Thr Tyr Trp Arg 120 Leu Glu Asn Ile Ala Lys Thr Glu Pro Pro Lys Ile Phe Arg Val Lys 130 135 Pro Val Leu Gly Ile Lys Arg Met Ile Gln Ile Glu Trp Ile Lys Pro 150 155 Glu Leu Ala Pro Val Ser Ser Asp Leu Lys Tyr Thr Leu Arg Phe Arg Thr Val Asn Ser Thr Ser Trp Met Glu Val Asn Phe Ala Lys Asn Arg 180 185 Lys Asp Lys Asn Gln Thr Tyr Asn Leu Thr Gly Leu Gln Pro Phe Thr Glu Tyr Val Ile Ala Leu Arg Cys Ala Val Lys Glu Ser Lys Phe Trp 215 Ser Asp Trp Ser Gln Glu Lys Met Gly Met Thr Glu Glu Gly Lys 225 230 235 240

Leu Leu Pro Ala Ile Pro Val Leu Ser Thr Leu Val 245 250

<210> 14

<211> 652

<212> PRT

<213> Homo sapiens

<400> 14

Met Lys Leu Ser Pro Gln Pro Ser Cys Val Asn Leu Gly Met Met Trp  $1 \hspace{1.5cm} 5 \hspace{1.5cm} 10 \hspace{1.5cm} 15$ 

Thr Trp Ala Leu Trp Met Leu Pro Ser Leu Cys Lys Phe Ser Leu Ala 20 25 30

Ala Leu Pro Ala Lys Pro Glu Asn Ile Ser Cys Val Tyr Tyr Tyr Arg 35 40 45

Lys Asn Leu Thr Cys Thr Trp Ser Pro Gly Lys Glu Thr Ser Tyr Thr 50 60

Gln Tyr Thr Val Lys Arg Thr Tyr Ala Phe Gly Glu Lys His Asp Asn 65 70 75 80

Cys Thr Thr Asn Ser Ser Thr Ser Glu Asn Arg Ala Ser Cys Ser Phe 85 90 95

Phe Leu Pro Arg Ile Thr Ile Pro Asp Asn Tyr Thr Ile Glu Val Glu
100 105 110

Ala Glu Asn Gly Asp Gly Val Ile Lys Ser His Met Thr Tyr Trp Arg 115 120 125

Leu Glu Asn Ile Ala Lys Thr Glu Pro Pro Lys Ile Phe Arg Val Lys 130 135 140

Pro Val Leu Gly Ile Lys Arg Met Ile Gln Ile Glu Trp Ile Lys Pro 145 150 155 160

Glu Leu Ala Pro Val Ser Ser Asp Leu Lys Tyr Thr Leu Arg Phe Arg 165 170 175

Thr Val Asn Ser Thr Ser Trp Met Glu Val Asn Phe Ala Lys Asn Arg 180 185 190

Lys Asp Lys Asn Gln Thr Tyr Asn Leu Thr Gly Leu Gln Pro Phe Thr 195 200 205

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Glu	Туг 210	Val	Ile	Ala	Leu	Arg 215	Cys	Ala	Val	Lys	Glu 220	Ser	Lys	Phe	Trp
Ser 225	Asp	Trp	Ser	Gln	Glu 230	Lys	Met	Gly	Met	Thr 235	Glu	Glu	Glu	Ala	Pro 240
Cys	Gly	Leu	Glu	Leu 245	Trp	Arg	Val	Leu	<b>Lys</b> 250	Pro	Ala	Glu	Ala	Asp 255	Gly
Arg	Arg	Pro	Val 260	Arg	Leu	Leu	Trp	Lys 265	Lys	Ala	Arg	Gly	Ala 270	Pro	Val
Leu	Glu	Lys 275	Thr	Leu	Gly	Tyr	Asn 280	Ile	Trp	Tyr	Tyr	Pro 285	Glu	Ser	Asn
Thr	Asn 290	Leu	Thr	Glu	Thr	Met 295	Asn	Thr	Thr	Asn	Gln 300	Gln	Leu	Glu	Leu
His 305	Leu	Gly	Gly	Glu	Ser 310	Phe	Trp	Val	Ser	Met 315	Ile	Ser	Tyr	Asn	Ser 320
Leu	Gly	Lys	Ser	Pro 325	Val	Ala	Thr	Leu	Arg 330	Ile	Pro	Ala	Ile	Gln 335	Glu
Lys	Ser	Phe	Gln 340	Cys	Ile	Glu	Val	Met 345	Gln	Ala	Сув	Val	Ala 350	Glu	Asp
Gln	Leu	Val 355	Val	Lys	Trp	Gln	Ser 360	Ser	Ala	Leu	Asp	Val 365	Asn	Thr	Trp
Met	Ile 370	Glu	Trp	Phe	Pro	Asp 375	Val	Asp	Ser	Glu	Pro 380	Thr	Thr	Leu	Ser
Trp 385	Glu	Ser	Val	Ser	Gln 390	Ala	Thr	Asn	Trp	Thr 395	Ile	Gln	Gln	Asp	Lys 400
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qzA	Lys	Val	Gly 420	Glu	Pro	Tyr	Ser	Ile 425	Gln	Ala	Tyr	Ala	Lys 430	Glu	Gly
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Phe Ser Lys Thr Val Asn Ser Ser Ile Leu Gln Tyr Gly Leu Glu Ser 485 490

Leu Lys Arg Lys Thr Ser Tyr Ile Val Gln Val Met Ala Asn Thr Ser 505

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Ser Val Phe Glu Ile Ile Leu Ile Thr Ser Leu Ile Gly Gly Leu 535

Leu Ile Leu Ile Leu Thr Val Ala Tyr Gly Leu Lys Lys Pro Asn

Lys Leu Thr His Leu Cys Trp Pro Thr Val Pro Asn Pro Ala Glu Ser 565 570

Ser Ile Ala Thr Trp His Gly Asp Asp Phe Lys Asp Lys Leu Asn Leu 585 580

Lys Glu Ser Asp Asp Ser Val Asn Thr Glu Asp Arg Ile Leu Lys Pro 600

Cys Ser Thr Pro Ser Asp Lys Leu Val Ile Asp Lys Leu Val Val Asn

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Leu Lys Arg Lys Thr Ser Tyr Ile Val Gln Val Met Ala Ser Thr Ser 500 505 510

Ala Gly Gly Thr Asn Gly Thr Ser Ile Asn Phe Lys Thr Leu Ser Phe 515 520 525

Ser Val Phe Glu Ile Ile Leu Ile Thr Ser Leu Ile Gly Gly Gly Leu 530 540

Leu Ile Leu Ile Leu Thr Val Ala Tyr Gly Leu Lys Lys Pro Asn 545 550 555 560

Lys Leu Thr His Leu Cys Trp Pro Thr Val Pro Asn Pro Ala Glu Ser 565 570 575

Lys Glu Ser Asp Asp Ser Val Asn Thr Glu Asp Arg Ile Leu Lys Pro 595 600 605

Cys Ser Thr Pro Ser Asp Lys Leu Val Ile Asp Lys Leu Val Val Asn 610 615 620

Phe Gly Asn Val Leu Gln Glu Ile Phe Thr Asp Glu Ala Arg Thr Gly 625 630 630 635

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Arg Ser His Asn Phe His Asn Leu Thr Thr Glu Glu Gly Pro Ser Thr

35 40 45

Pro Ile Gly Thr Leu Lys Pro Gly Leu Val Ile Lys Ala Val Arg Lys 50 60

Leu Leu Met Asn Asp Ser Asp Gln Gly Gly Lys Leu Thr Thr Gly Val 65 70 75 80

Phe Thr Pro Gln Gln Leu Ala Asn Thr Thr Asn Gln Gly Leu Ser Arg 85 90 95

Cys Leu Ser Arg Phe Lys Lys Val Ile Arg Ala Met Leu Met Met Lys 100 105 110

Ile Lys Leu Lys Arg Ile Thr Asn Ile Asn Cys Ser Gly His Ile Trp \$115\$ \$120\$

Val Glu Pro Ala Thr Ile Phe Lys Met Gly Met Asn Ile Ser Ile Tyr 130 135 140

Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu His Phe Tyr 145 150 155 160

Lys Asn Gly Ile Lys Glu Arg Phe Gln Ile Thr Arg Ile Asn Lys Thr 165 170 175

Thr Ala Arg Leu Trp Tyr Lys Asn Phe Leu Glu Pro His Ala Ser Met 180 185 190

Tyr Cys Thr Ala Glu Cys Pro Lys His Phe Gln Glu Thr Leu Ile Cys 195 200 205

Gly Lys Asp Ile Ser Ser Gly Phe Cys Ile Thr Asp Tyr Ser Gln Lys 210 215 220

Pro Ser Gln Val Leu Ala Gly Gly Pro Leu Ser Pro Asn Pro Thr Pro 225 230 235 240

Gly Asn Val Glu Asp Pro Pro Asp Ile Pro Asp Glu Val Thr Cys Val 245 250 255

Ile Tyr Glu Tyr Ser Gly Asn Met Thr Cys Thr Trp Asn Ala Gly Lys 260 265 270

Leu Thr Tyr Ile Asp Thr Lys Tyr Val Val His Val Lys Ser Leu Glu 275 280 285

35

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Ser Ile Tyr Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu

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<sup>&</sup>lt;213> Homo sapiens

<sup>&</sup>lt;400> 21

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His	Leu	Ser	Asn 500	Asn	Asn	Glu	Ile	Thr 505	Ser	Leu	Thr	Leu	Lys 510	Pro	Pro
Val	Asp	Ser 515	Leu	Asp	Ser	Gly	Asn 520	Asn	Pro	Arg	Leu	Gln 525	Lys	His	Pro
Asn	Phe 530	Ala	Phe	Ser	Val	Ser 535	Ser	Val	Asn	Ser	Leu 540	Ser	Asn	Thr	Ile
Phe	Leu	Gly	Glu	Leu	Ser	Leu	Ile	Leu	Asn	Gln	Gly	Glu	Cys	Ser	Ser

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960

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His Ile Trp Val Glu Pro Ala Thr Ile Phe Lys Met Gly Met Asn Ile 35 - 40

Ser Ile Tyr Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu 50 55

His Phe Tyr Lys Asn Gly Ile Lys Glu Arg Phe Gln Ile Thr Arg Ile

Asn Lys Thr Thr Ala Arg Leu Trp Tyr Lys Asn Phe Leu Glu Pro His

Ala Ser Met Tyr Cys Thr Ala Glu Cys Pro Lys His Phe Gln Glu Thr 100 105

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Thr 145	Trp	Asn	Ala	Gly	Lys 150	Leu	Thr	Tyr	Ile	Asp 155	Thr	Lys	Tyr	Val	Val 160
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Glu Leu Met Asn Asn Ser Ser Glu Gln Val Leu Tyr Val Asp Pro 360

Met Ile Thr Glu Ile Lys Glu Ile Phe Ile Pro Glu His Lys Pro Thr 375 380

Asp Tyr Lys Lys Glu Asn Thr Gly Pro Leu Glu Thr Arg Asp Tyr Pro 385

Gln Asn Ser Leu Phe Asp Asn Thr Thr Val Val Tyr Ile Pro Asp Leu 405 410

Asn Thr Gly Tyr Lys Pro Gln Ile Ser Asn Phe Leu Pro Glu Gly Ser

His Leu Ser Asn Asn Glu Ile Thr Ser Leu Thr Leu Lys Pro Pro 440 435

Val Asp Ser Leu Asp Ser Gly Asn Asn Pro Arg Leu Gln Lys His Pro 455

Asn Phe Ala Phe Ser Val Ser Ser Val Asn Ser Leu Ser Asn Thr Ile 470 , 475

Phe Leu Gly Glu Leu Ser Leu Ile Leu Asn Gln Gly Glu Cys Ser Ser 485 490 495

Pro Asp Ile Gln Asn Ser Val Glu Glu Glu Thr Thr Met Leu Glu 500 505

Asn Asp Ser Pro Ser Glu Thr Ile Pro Glu Gln Thr Leu Leu Pro Asp 515 520 525

Glu Phe Val Ser Cys Leu Gly Ile Val Asn Glu Glu Leu Pro Ser Ile 530 535 540

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Leu Phe Ser Trp Cys His Gly Gly Ile Thr Asn Ile Asn Cys Ser Gly

His Ile Trp Val Glu Pro Ala Thr Ile Phe Lys Met Gly Met Asn Ile

Ser Ile Tyr Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu 50 55 60

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His Phe Tyr Lys Asn Gly Ile Lys Glu Arg Phe Gln Ile Thr Arg Ile Asn Lys Thr Thr Ala Arg Leu Trp Tyr Lys Asn Phe Leu Glu Pro His Ala Ser Met Tyr Cys Thr Ala Glu Cys Pro Lys His Phe Gln Glu Thr Leu Ile Cys Gly Lys Asp Ile Ser Ser Gly Tyr Pro Pro Asp Ile Pro Asp Glu Val Thr Cys Val Ile Tyr Glu Tyr Ser Gly Asn Met Thr Cys Thr Trp Asn Ala Gly Lys Leu Thr Tyr Ile Asp Thr Lys Tyr Val Val His Val Lys Ser Leu Glu Thr Glu Glu Glu Gln Tyr Leu Thr Ser Ser Tyr Ile Asn Ile Ser Thr Asp Ser Leu Gln Gly Gly Lys Lys Tyr Leu Val Trp Val Gln Ala Ala Asn Ala Leu Gly Met Glu Glu Ser Lys Gln Leu Gln Ile His Leu Asp Asp Ile Val Ile Pro Ser Ala Ala Val Ile Ser Arg Ala Glu Thr Ile Asn Ala Thr Val Pro Lys Thr Ile Ile Tyr Trp Asp Ser Gln Thr Thr Ile Glu Lys Val Ser Cys Glu Met Arg Tyr Lys Ala Thr Thr Asn Gln Thr Trp Asn Val Lys Glu Phe Asp Thr Asn Phe Thr Tyr Val Gln Gln Ser Glu Phe Tyr Leu Glu Pro Asn Ile Lys Tyr Val Phe Gln Val Arg Cys Gln Glu Thr Gly Lys Arg Tyr Trp Gln Pro Trp Ser Ser Leu Phe Phe His Lys Thr Pro Glu Thr Val Pro

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Gln Val Thr Ser Lys Ala Phe Gln His Asp Thr Trp Asn Ser Gly Leu . 325 330 335

Thr Val Ala Ser Ile Ser Thr Gly His Leu Thr Ser Gly Leu Lys Glu

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His Ile Trp Val Glu Pro Ala Thr Ile Phe Lys Met Gly Met Asn Ile 35 40 45

Ser Ile Tyr Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu 55

His Phe Tyr Lys Asn Gly Ile Lys Glu Arg Phe Gln Ile Thr Arg Ile 70

Asn Lys Thr Thr Ala Arg Leu Trp Tyr Lys Asn Phe Leu Glu Pro His 85 90

Ala Ser Met Tyr Cys Thr Ala Glu Cys Pro Lys His Phe Gln Glu Thr 100

Leu Ile Cys Gly Lys Asp Ile Ser Ser Gly Tyr Pro Pro Asp Ile Pro

Asp Glu Val Thr Cys Val Ile Tyr Glu Tyr Ser Gly Asn Met Thr Cys 130 135

Thr Trp Asn Ala Gly Lys Leu Thr Tyr Ile Asp Thr Lys Tyr Val Val 145 150 155 160

48

His Val Lys Ser Leu Glu Thr Glu Glu Glu Gln Tyr Leu Thr Ser

Ser Tyr Ile Asn Ile Ser Thr Asp Ser Leu Gln Gly Gly Lys Lys Tyr 180 185 190

Leu Val Trp Val Gln Ala Ala Asn Ala Leu Gly Met Glu Glu Ser Lys 195 200

Gln Leu Gln Ile His Leu Asp Asp Ile Val Ile Pro Ser Ala Ala Val 210 215

Ile Ser Arg Ala Glu Thr Ile Asn Ala Thr Val Pro Lys Thr Ile Ile 230

Tyr Trp Asp Ser Gln Thr Thr Ile Glu Lys Val Ser Cys Glu Met Arg 245 250

Tyr Lys Ala Thr Thr Asn Gln Thr Trp Asn Val Lys Glu Phe Asp Thr

Asn Phe Thr Tyr Val Gln Gln Ser Glu Phe Tyr Leu Glu Pro Asn Ile

Lys Tyr Val Phe Gln Val Arg Cys Gln Glu Thr Gly Lys Arg Tyr Trp 290 295 300

Gln Pro Trp Ser Ser Pro Phe Phe His Lys Thr Pro Glu Thr Val Pro 310 315

Gln Val Thr Ser Lys Ala Phe Gln His Asp Thr Trp Asn Ser Gly Leu 330

Thr Val Ala Ser Ile Ser Thr Gly His Leu Thr Ser Asp Asn Arg Gly 345

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Tyr Phe Tyr Lys Asn Gly Phe Lys Glu Glu Phe Asp Ile Thr Arg Ile 65 70 75 80

Asn Arg Thr Thr Ala Arg Ile Trp Tyr Lys Gly Phe Ser Glu Pro His
85 90 95

Ala Tyr Met His Cys Thr Ala Glu Cys Pro Gly His Phe Gln Glu Thr 100 105 110

Leu Ile Cys Gly Lys Asp Ile Ser Ser Gly His Pro Pro Asp Ala Pro 115 120 125

Ser Asn Leu Thr Cys Val Ile Tyr Glu Tyr Ser Gly Asn Met Thr Cys 130 140

Thr Trp Asn Thr Gly Lys Pro Thr Tyr Ile Asp Thr Lys Tyr Ile Val 145 150 155 160

His Val Lys Ser Leu Glu Thr Glu Glu Glu Glu Gln Tyr Leu Ala Ser 165 170 175

Ser Tyr Val Lys Ile Ser Thr Asp Ser Leu Gln Gly Ser Arg Lys Tyr 180 185 190

Leu Val Trp Val Gln Ala Val Asn Ser Leu Gly Met Glu Asn Ser Gln
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Gln Leu His Val His Leu Asp Asp Ile Val Ile Pro Ser Ala Ser Ile 210 215 220

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50

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Arg Asp Cys Pro Leu Gly Met Leu Ser Thr Ser Ser Ser Val Val Tyr 490 Ile Pro Asp Leu Asn Thr Gly Tyr Lys Pro Gln Val Ser Asn Val Pro 500 505 Pro Gly Gly Asn Leu Phe Ile Asn Arg Asp Glu Arg Asp Pro Thr Ser 515 520 525 Leu Glu Thr Thr Asp Asp His Phe Ala Arg Leu Lys Thr Tyr Pro Asn 530 Phe Gln Phe Ser Ala Ser Ser Met Ala Leu Leu Asn Lys Thr Leu Ile 545 560 550 555 Leu Asp Glu Leu Cys Leu Val Leu Asn Gln Gly Glu Phe Asn Ser Leu Asp Ile Lys Asn Ser Arg Gln Glu Glu Thr Ser Ile Val Leu Gln Ser 585 Asp Ser Pro Ser Glu Thr Ile Pro Ala Gln Thr Leu Leu Ser Asp Glu 595 600 605 Phe Val Ser Cys Leu Ala Ile Gly Asn Glu Asp Leu Pro Ser Ile Asn Ser Tyr Phe Pro Gln Asn Val Leu Glu Ser His Phe Ser Arg Ile Ser 625 630 635 Leu Phe Gln Lys <210> 28 <211> 2181 <212> DNA <213> Mus musculus <400> 28 atgaagcctc tgggtgtgaa cgctggaata atgtggacct tggcactgtg ggcattctct 60 ttcctctgca aattcagcct ggcagtcctg ccgactaagc cagagaacat ttcctgcgtc 120 ttttacttcg acagaaatct gacttgcact tggagaccag agaaggaaac caatgatacc 180 agctacattg tgactttgac ttactcctat ggaaaaagca attatagtga caatgctaca 240 gaggetteat attetttee cegtteetgt geaatgeece cagacatetg cagtgttgaa 300

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ccatctatta	attcttactt	tccacagaac	gttttggaaa	gccatttcag	tagaatttca	1920
ctcttccaaa	agtag					1935

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#### **PCT**

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- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

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(54) Title: HEMATOPOIETIN RECEPTORS HPR1 AND HPR2

(57) Abstract: This invention relates to human and murine HPR1 and human and murine HPR2 polypeptides, new members of the hematopoietin receptor polypeptide family; to methods of making such HPR1 and HPR2 polypeptides; to non-human mammals in which the endogenous genomic sequences encoding HPR1 and/or HPR2 polypeptides have been partially or completely inactivated; to methods of using HPR1 or HPR2 polypeptides to identify compounds that alter HPR1 or HPR2 polypeptide activities; and to methods of preparing medicaments for and/or treating conditions associated with hematopoietin receptor function.

Internation No PCT/US 01/31634

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C12N15/12 C07K14/705 C07K14/715

According to International Patent Classification (IPC) or to both national classification and IPC

#### **B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)  $I\,PC\,\,7\,\,\,\,C07\,K$ 

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

WPI Data, PAJ, EPO-Internal, MEDLINE, CHEM ABS Data, BIOSIS, SEQUENCE SEARCH, EMB

C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of	the relevant passages	Relevant to claim No.
Y	WO 98 31811 A (DONALDSON DEBRINST (US); COLLINS MARY (US); 23 July 1998 (1998-07-23) the whole document	RA D ;GENETICS NEBEN TA)	1,5-7, 12-35
Υ	LARSEN A ET AL: "EXPRESSION HUMAN GRANULOCYTE COLONY-STIM FACTORRECEPTOR: A STRUCTURAL HEMATOPOIETIN RECEPTOR, IMMUN AND FIBRONECTIN DOMAINS" JOURNAL OF EXPERIMENTAL MEDICULARY JOURNAL OF EXPERIMENTAL MEDICULARY STATES AND STATES A	NULATING MOSAIC OF NOGLOBULIN, CINE, TOKYO,	1,5-7, 12-35
X Furt	her documents are listed in the continuation of box C.	Patent family members are listed	in annex.
"A" docume consic "E" earlier of filing of "L" docume which citatio "O" docume other "P" docume	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another n or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but han the priority date claimed	"T" later document published after the integration or priority date and not in conflict with cited to understand the principle or the invention  "X" document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the document of particular relevance; the cannot be considered to involve an indocument is combined with one or ments, such combination being obvious in the art.  "&" document member of the same patent	the application but eory underlying the slaimed invention to be considered to coument is taken alone claimed invention ventive step when the ore other such docutes to a person skilled family
	actual completion of the international search	Date of mailing of the international sea	arch report
	August 2002	1 3. 11. 02	
Name and I	mailing address of the ISA	Authorized officer	
	European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Armandola, E	

International Application No
PCT/VS 01/31634

C.(Continue	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	101/03 01/31001	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
Ρ,Υ	DATABASE EMBL [Online] SEQ. ID. NO: 17, 26 March 2001 (2001-03-26) MAEDA M. AND YAGUCHI N.: "Human haematopoietin receptor protein NR10.3" Database accession no. AAB51244 XP002209359 the whole document	1,5-7, 12-35	
P,X	-& WO 00 75314 A (CHUGAI RES INST FOR MOLECULAR ;MAEDA MASATSUGU (JP); YAGUCHI NORIK) 14 December 2000 (2000-12-14)	1,5-7, 12-35	
Е	WO 02 00721 A (ZYMOGENETICS INC) 3 January 2002 (2002-01-03) SEQ. ID. NO: 1 and SEQ. ID. NO: 46	1,5-7, 12-35	
L	DATABASE EMBL [Online] Sequence 1 from patent W00200721, 15 February 2002 (2002-02-15) SPRECHER C.A. ET AL.: "Cytokine receptor zcytor17" Database accession no. AX365149 XP002209360		
L	the whole document -& DATABASE EMBL [Online] Human zcytor17 protein sequence SEQ.ID.NO:46, 1 May 2002 (2002-05-01) SPRECHER C.A. ET AL.: "Isolated polynucleotide encoding a cytokine receptor zcytor17." Database accession no. AAB05730 XP002209361 L document cited to provide information on the relevant sequence disclosed in W00200721 the whole document		
E	WO 01 93983 A (STEPHAN JEAN PHILIPPE F; BAKER KEVIN P (US); DESNOYERS LUC (US); G) 13 December 2001 (2001-12-13) SEQ. ID. NO: 91 and NO: 92	1,5-7, 12-35	

International Application No
PC1/50 01/31634

	ation) DOCUMENTS CONSIDERED TO BE RELEVANT  Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
Category °	Citation of document, with moleculon, where appropriate, of the relevant passages		
L	DATABASE EMBL [Online] Sequence 91 from patent W00193983, 13 February 2002 (2002-02-13) BAKER K.P. ET AL.: "Secreted transmembrane polypeptides and nucleic acids encoding the same." Database accession no. AX358838 XP002209362	•	
L	the whole document -& DATABASE EMBL [Online] Sequence 92 from Patent W00193983, 13 February 2002 (2002-02-13) BAKER K.P. ET AL.: "Secreted and transmembrane polypeptides and nucleic acids encoding the same" Database accession no. AX358839 XP002209363 L document cited to provide information on the relevant sequence disclosed in W00193983 the whole document		

ional application No. PCT/US 01/31634

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)			
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:			
1. X Claims Nos.: 28,30-35 because they relate to subject matter not required to be searched by this Authority, namely:  Although claims 28 and 30-35 are directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged			
effects of the compound/composition.  2. Claims Nos.: 5,12-15,18, 27-30, 32-35 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:  see FURTHER INFORMATION sheet PCT/ISA/210			
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).			
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)			
This International Searching Authority found multiple inventions in this international application, as follows:			
see additional sheet			
1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.			
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.			
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:			
No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:  1,5,12-35(partially), 6,7 (completely)			
Remark on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.			

### FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1, 5, 12-35 (partially), 6,7 (completely)

An isolated polypeptide HPR1 comprising the amino acid sequence of SEQ. ID. NO: 4, subsequences thereof, fusions of subsequences of SEQ. ID. NO: 4, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequences SEQ. ID. NO: 3, SEQ. ID. NO: 5 and subsequences thereof, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

2. Claims: 1.5.12-35 (all partially)

HPR1 amino acid sequences comprising fragments of SEQ. ID. NO: 4 fused to fragments of SEQ. ID. NO: 1, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

3. Claims: 1,5,12-35 (all partially)

HPR1 amino acid sequences comprising fragments of SEQ. ID. NO: 4 fused to fragments of SEQ. ID. NO: 2, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

4. Claims: 1.5,12-35 (all partially)

HPR1 amino acid sequences comprising fragments of SEQ. ID. NO: 4 fused to fragments of SEQ. ID. NO: 10, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

5. Claims: 1,5,12-35 (all partially)

HPR1 amino acid sequences comprising fragments of SEQ. ID. NO: 4 fused to fragments of SEQ. ID. NO: 11, isolated polynucleotides encoding the polypeptides, expression

### FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

6. Claims: 2 and 8 (completely), 5 and 12-35 (partially)

An isolated polypeptide HPR1 comprising the amino acid sequence of SEQ. ID. NO: 12, subsequences thereof, fusions of subsequences of SEQ. ID. NO: 12, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequence SEQ. ID. NO: 28, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

7. Claims: 3, 5,9, 12-35 (all partially)

An isolated polypeptide HPR2 comprising the amino acid sequence SEQ. ID. NO: 23, subsequences thereof, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequence SEQ. ID. NO: 22, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

8. Claims: 3, 5,9, 12-35 (all partially)

An isolated polypeptide HPR2 comprising the amino acid sequence SEQ. ID. NO: 25, subsequences thereof, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequence SEQ. ID. NO: 24, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

9. Claims: 3, 5,12-35 (all partially)

HPR2 amino acid sequences comprising fragments of SEQ. ID. NO: 23 fused to SEQ. ID. NO: 16 or to fragments of SEQ. ID. NO: 16 or to SEQ. ID. NO: 17 or to SEQ. ID. NO: 18, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

#### FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

10. Claims: 4, 11 (completely), 5 and 12-35 (partially)

An isolated polypeptide HPR2 comprising the amino acid sequence SEQ. ID. NO: 27, subsequences thereof, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequence SEQ. ID. NO: 29, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

11. Claims: 5, 9 and 12-35 (all partially)

Polynucleotide sequences comprising SEQ. ID. NO: 19 or SEQ. ID. NO: 20, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

# FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 5,12-15,18, 27-30, 32-35

Present claims 5,12-15,18, 27-30, 32-35 relate to an extremely large number of possible compounds/products. Support within the meaning of Article 6 PCT and/or disclosure within the meaning of Article 5 PCT is to be found, however, for only a very small proportion of the compounds/products. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible. Consequently, the search has been carried out for those parts of the claims which appear to be supported and disclosed, namely those parts relating to the DNAs or proteins for which a sequence listing has been provided and to antibodies recognizing the claimed proteins.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

# INTERNATIONAL SEARCH REPORT

rmation on patent family members

Internation No
PCT/US 01/31634

	•						
c	Patent document ited in search report		Publication date		Patent family member(s)		Publication date
V	vO 9831811	A	23-07-1998	AU EP JP WO	5733898 1005552 2001508309 9831811	A1 T	07-08-1998 07-06-2000 26-06-2001 23-07-1998
V	√O 0075314	Α	14-12-2000	AU EP WO	4952400 1188830 0075314	A1	28-12-2000 20-03-2002 14-12-2000
V	√0 0200721	Α	03-01-2002	AU WO	7303201 0200721		08-01-2002 03-01-2002
	√O 0193983	A	13-12-2001	- USOUUUUUPPSOOOOOSSSSSSSSSSSSSSSSSSSSSSS	2002106743 2002052027 2055401 2590901 5460100 6501901 6531101 6802801 1250426 1240325 2002119130 0140466 0146420 0168848 0192331 0193983 2002160392 2002098505 2002098505 2002098507 2002090681 2002127584 2002127584 2002146709 2002132240 2002132240 2002072496 2002142961 2002072067 2002072067 2002072067 2002072067 2002132253 2002132252 2002127576 2002132253 2002132253 2002132253 2002132253 2002132253 200213253 200213253 200213253 200213253 200213253 200213253 200213253 200213253 200213253 200213253 200213253	-A11 -A11 -A11 -A11 -A11 -A11 -A11 -A11	08-08-2002 02-05-2002 12-06-2001 03-07-2001 18-12-2000 11-12-2001 17-12-2001 24-09-2001 23-10-2002 18-09-2001 28-06-2001 28-06-2001 28-06-2001 20-09-2001 13-12-2002 11-07-2002 25-07-2002 25-07-2002 25-07-2002 11-07-2002 12-09-2002 11-07-2002 12-09-2002 13-06-2001 25-09-2002 25-09-2002 212-09-2002 212-09-2002 212-09-2001 26-03-2001 26-03-2001 26-03-2001 26-03-2001 26-03-2001 28-03-2001 08-03-2001

# INTERNATIONAL SEARCH REPORT

rmation on patent family members

International Application No
PCT/OS 01/31634

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 0193983	A	WO 0140465 A2 WO 0166740 A2 US 2002106728 A1 AU 7197301 A	07-06-2001 13-09-2001 08-08-2002 05-02-2002

# **CORRECTED VERSION**

# (19) World Intellectual Property Organization

International Bureau



# - 1 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1881 | 1

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(21) International Application Number: PCT/US01/31634

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(25) Filing Language: English

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- (71) Applicant (for all designated States except US): IM-MUNEX CORPORATION [US/US]; Law Department, 51 University Street, Seattle, WA 98101 (US).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): COSMAN, David, J. [GB/US]; 51 University Street, Seattle, WA 98101 (US). MOSLEY, Bruce, A. [US/US]; 51 University Street, Seattle, WA 98101 (US). BIRD, Timothy, A. [GB/US]; 51 University Street, Seattle, WA 98101 (US). DUBOSE, Robert, F. [US/US]; 51 University Street, Seattle, WA 98101 (US). WILEY, Steven, R. [US/US]; 51 University Street, Seattle, WA 98101 (US).
- (74) Agent: SPRUNGER, Suzanne, A.; Immunex Corporation, Law Department, 51 University Street, Seattle, WA 98101 (US).

- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

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17 July 2003

(15) Information about Correction: see PCT Gazette No. 29/2003 of 17 July 2003, Section II

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.



(57) Abstract: This invention relates to human and murine HPR1 and human and murine HPR2 polypeptides, new members of the hematopoietin receptor polypeptide family; to methods of making such HPR1 and HPR2 polypeptides; to non-human mammals in which the endogenous genomic sequences encoding HPR1 and/or HPR2 polypeptides have been partially or completely inactivated; to methods of using HPR1 or HPR2 polypeptides to identify compounds that alter HPR1 or HPR2 polypeptide activities; and to methods of preparing medicaments for and/or treating conditions associated with hematopoietin receptor function.





# HEMATOPOIETIN RECEPTORS HPR1 AND HPR2

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This application claims the benefit under 35 U.S.C. 119(e) of U.S. provisional applications Serial No. 60/238,706, filed 06 October 2000; Serial No. 60/240,476, filed 13 October 2000; and Serial No. 60/270,282, filed 20 February 2001; all of which are incorporated by reference herein.

## **FIELD OF THE INVENTION**

This invention relates to new human and murine hematopoietin receptor polypeptides HPR1 and HPR2, and to methods of making and using HPR1 and HPR2 polypeptides.

### **BACKGROUND OF THE INVENTION**

The hematopoietin receptor polypeptides are a related group of Type I membrane protein receptors, and in some cases soluble forms of those receptors; this family of polypeptides has variously been called the cytokine receptor family or the hematopoietin receptor family. There are other families of receptors that bind cytokines or growth factors, such as the IL-1 receptor family, the TNF receptor family, and the EGF receptor family, but the hematopoietin receptor family is considered to be a distinct group or family of receptors based on certain characteristic structural features or motifs that are shared by members of this family. Some of the members of the hematopoietin receptor family are gp130, the granulocyte colony-stimulating factor receptor (GCSFR), leukemia inhibitory factor receptor (LIF-R), the alpha chains and the common beta chain of the IL-3 and IL-5 receptors, etc.; the hematopoietin receptor family contains more than 20 different polypeptides.

Common structural features of the hematopoietin receptor family of polypeptides include at least one extracellular cytokine receptor domain, which usually contains four cysteines and a WSXWS motif (where W is tryptophan, S is serine, and X indicates any amino acid), and, in most members of the family, a transmembrane and a cytoplasmic domain. The extracellular cytokine receptor domain is involved in ligand-binding activity, and the intracellular domain of a 'signaling' subfamily of hematopoietin receptors has a signal transduction function, transmitting the signal generated by ligand binding to a signal transduction pathway that results in the expression of genes involved in cell proliferation, differentiation, and/or activation. These activities of the hematopoietin receptor polypeptide family are mediated through interactions with cytokine ligands and other ligand-binding receptor molecules, with ligand binding to the cytokine receptor domain of hematopoietin receptor polypeptides and facilitating homo- or heterotypic interactions between receptor polypeptides, bringing the cytoplasmic domains of receptors into proximity with each other. Many of the cytokine ligands (such as IL-2, IL-6, or ciliary neurotrophic factor or CNTF, for example) interact with more than one type of heteromeric hematopoietin receptor complex, often with differing affinities, and "common" hematopoietin receptor polypeptides such as gp130 are involved in several different heteromeric receptor complexes that bind a variety of ligands. Because of their ligand-binding and intracellular signaling activities, hematopoietin receptor polypeptides are associated with a wide variety of conditions involving cytokine-influenced cell proliferation, differentiation, or activation. For example, interaction of the gp130 hematopoietin receptor polypeptide with its binding partners is involved in the

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normal upregulation of cardiac myocyte proliferation ("hypertrophy") in response to biomechanical stress on the heart, as lack of gp130 leads to heart failure under those conditions (Hirota *et al.*, 1999, *Cell* 97(2): 189-198). Hematopoietin receptors are also involved in the activation or stimulation of cells in response to environmental factors, for example the activation of hepatocytes in the acute-phase inflammatory response to injury (Taga and Kishimoto, 1992, *Crit Rev Immunol.* 11(5): 265-280; Neben and Turner, 1993, *Stem Cells* 11 Suppl 2: 156-162). Hematopoietin receptor family polypeptides generally are constitutively expressed in many different cell types throughout development, but the expression levels of hematopoietin receptor polypeptides may be up- or downregulated in response to stimuli, and some members of the family exhibit more restricted patterns of expression in particular tissues. Characteristics and activities of the hematopoietin receptor polypeptide family are described further in the following references, which are incorporated by reference herein: Drachman and Kaushansky, 1995, Curr Opin Hematol. 2(1): 22-28; Ihle, 1995, Nature 377(6550): 591-594; Taga and Kishimoto, 1995, Curr Opin Immunol. 7(1): 17-23; Ihle *et al.*, 1995, Annu Rev Immunol. 13: 369-398; Theze, 1994, Eur Cytokine Netw. 5(4): 353-368; Ihle et al., 1994, Trends Biochem Sci. 19(5): 222-227; Cosman, 1993, Cytokine 5(2): 95-106; and Onishi *et al.*, 1998, Int Rev Immunol. 16(5-6): 617-634.

In order to develop more effective treatments for disorders such as neurological, cardiac, hematopoietic, immunological, hepatic, and pulmonary conditions and diseases involving cell proliferation, differentiation, or activation, including neoplastic transformation or proliferation of virus-infected or cancerous cells, information is needed about previously unidentified members of the hematopoietin receptor polypeptide family.

# **SUMMARY OF THE INVENTION**

The present invention is based upon the discovery of new human hematopoietin receptor family members, HPR1 and HPR2.

The invention provides an isolated polypeptide consisting of, consisting essentially of, or more preferably, comprising an amino acid sequence selected from the group consisting of:

- (a) the amino acid sequence of SEQ ID NO:4;
- (b) amino acids 56 through 77 of SEQ ID NO:1;
- an amino acid sequence selected from the group consisting of: amino acids 1 through 55 of SEQ ID NO:1; amino acids 5 through 40 of SEQ ID NO:2; amino acids 1 through 32 of SEQ ID NO:4; amino acids 1 through 525 of SEQ ID NO:4; amino acids 20 through 32 of SEQ ID NO:4; amino acids 33 through 134 of SEQ ID NO:4; amino acids Xaa1 through Xaa2 of SEQ ID NO:4, wherein Xaa1 is selected from the group consisting of amino acids 33 through 43 of SEQ ID NO:4 and Xaa2 is selected from the group consisting of amino acids 228 through 241 of SEQ ID NO:4; amino acids 33 through 238 of SEQ ID NO:4; amino acids 33 through 525 of SEQ ID NO:4; amino acids 33 through 525 of SEQ ID NO:4; amino acids 33 through 745 of SEQ ID NO:4; amino acids 44 through 94 of SEQ ID NO:4; amino acids 139 through 241 of SEQ ID NO:4; amino acids 242 through 326 of SEQ ID NO:4; amino acids 242 through 514 of SEQ ID NO:4; amino acids 337 through 419 of SEQ ID NO:4; amino acids 242 through 514 of SEQ ID NO:4; amino acids 337 through 419 of SEQ ID NO:4;

amino acids 433 through 514 of SEQ ID NO:4; amino acids 526 through 556 of SEQ ID NO:4; amino acids 533 through 552 of SEQ ID NO:4; amino acids 553 through 745 of SEQ ID NO:4; amino acids 557 through 745 of SEQ ID NO:4; amino acids 563 through 573 of SEQ ID NO:4; amino acids 563 through 641 of SEQ ID NO:4; amino acids 567 through 581 of SEQ ID NO:4; amino acids 588 through 639 of SEQ ID NO:4; and amino acids 631 through 641 of SEQ ID NO:4;

- (d) fragments of the amino acid sequences of any of (a)-(c) comprising at least 20 contiguous amino acids;
- (e) fragments of the amino acid sequences of any of (a)-(c) comprising at least 30 contiguous amino acids;
- (f) fragments of the amino acid sequences of any of (a)-(c) having HPR1 polypeptide activity;
- (g) fragments of the amino acid sequences of any of (a)-(c) comprising cytokine receptor domain amino acid sequences;
  - (h) an allelic variant of any of (a)-(c);

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- (i) amino acid sequences comprising at least 20 amino acids and sharing amino acid identity with the amino acid sequences of any of (a)-(h), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99%, and at least 99.5%;
- an amino acid sequence of any of (a)-(i) wherein the polypeptide comprising said amino acid sequence also comprises an amino acid sequence selected from the group consisting of SEQ ID NO:10, SEQ ID NO:11, amino acids 652 though 745 of SEQ ID NO:4, a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least 20 contiguous amino acids; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least 30 contiguous amino acids; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 that is at least 25% of the length of the sequence of amino acids 652 though 745 of SEQ ID NO:4; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 that is at least 50% of the length of the sequence of amino acids 652 though 745 of SEQ ID NO:4; and a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4; and a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least one tyrosine residue;
- (k) an amino acid sequence of any of (a)-(j) wherein the polypeptide comprising said amino acid sequence does not comprise an amino acid sequence selected from the group consisting of amino acids 239 through 252 of SEQ ID NO:13; amino acids 643 through 652 of SEQ ID NO:14; and amino acids 652 through 662 of SEQ ID NO:15;
- (l) an amino acid sequence of (i)-(k), wherein a polypeptide comprising said amino acid sequence of (i)-(k) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (a)-(h); and
  - (m) an amino acid sequence of (i)-(l) having HPR1 polypeptide activity.

- 3 -

Preferably, such polypeptides are isolated HPR1 polypeptides or isolated polypeptides having HPR1 polypeptide activity.

Other aspects of the invention are isolated nucleic acids encoding polypeptides of the invention, with a preferred embodiment being an isolated nucleic acid consisting of, consisting essentially of, or more preferably, comprising a nucleotide sequence selected from the group consisting of:

(a) SEQ ID NO:3;

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- (b) SEQ ID NO:5;
- (c) nucleotides 132 through 2366 of SEQ ID NO:3; and
- (d) allelic variants of (a)-(c).

An additional preferred embodiment of the invention is an isolated nucleic acid consisting of, consisting essentially of, or more preferably, comprising a nucleotide sequence selected from the group consisting of nucleotides 1 through 137 of SEQ ID NO:3, nucleotides 138 through 228 of SEQ ID NO:3, nucleotides 229 through 346 of SEQ ID NO:3, nucleotides 347 through 528 of SEQ ID NO:3, nucleotides 529 through 680 of SEQ ID NO:3, nucleotides 681 through 846 of SEQ ID NO:3, nucleotides 847 through 926 of SEQ ID NO:3, nucleotides 927 through 1143 of SEQ ID NO:3, nucleotides 1144 through 1326 of SEQ ID NO:3, nucleotides 1327 through 1428 of SEQ ID NO:3, nucleotides 1429 through 1575 of SEQ ID NO:3, nucleotides 1576 through 1716 of SEQ ID NO:3, nucleotides 1717 through 1810 of SEQ ID NO:3, nucleotides 1811 through 1892 of SEQ ID NO:3, and nucleotides 1893 through 2480 of SEQ ID NO:3.

The invention provides an isolated polypeptide consisting of, consisting essentially of, or more preferably, comprising an amino acid sequence selected from the group consisting of:

- (a) the amino acid sequence of SEQ ID NO:21;
- (b) an amino acid sequence selected from the group consisting of: amino acids 1 through 177 of SEQ ID NO:16; amino acids 216 through 245 of SEQ ID NO:16; SEQ ID NO:17; SEQ ID NO:18; and amino acids 349 through 356 of SEQ ID NO:23;
- an amino acid sequence selected from the group consisting of: amino acids 1 through 23 of SEQ ID NO:21; amino acids 1 through 124 of SEQ ID NO:21; amino acids 1 through 318 of SEQ ID NO:21; amino acids 1 through 331 of SEQ ID NO:21; amino acids 1 through 355 of SEQ ID NO:21; amino acids Xaa1 through Xaa2 of SEQ ID NO:21, wherein Xaa1 is selected from the group consisting of amino acids 24 through 30 of SEQ ID NO:21 and Xaa2 is selected from the group consisting of amino acids 115 through 124 of SEQ ID NO:21; amino acids 24 through 331 of SEQ ID NO:21; amino acids 24 through 355 of SEQ ID NO:21; amino acids Xaa3 through Xaa4 of SEQ ID NO:21, wherein Xaa3 is selected from the group consisting of amino acids 125 through 133 of SEQ ID NO:21; amino acids 125 through 219 of SEQ ID NO:21; amino acids 309 through 331 of SEQ ID NO:21; amino acids 133 through 309 of SEQ ID NO:21; amino acids 224 through 320 of SEQ ID NO:21; amino acids 224 through 331 of SEQ ID NO:21; amino acids 224 through 331 of SEQ ID NO:21; amino acids 224 through 331 of SEQ ID NO:21; amino acids 224 through 331 of SEQ ID NO:21; amino acids 224 through 331 of SEQ ID NO:21; amino acids 234 through 331 of SEQ ID NO:21; amino acids 319 through 565 of SEQ ID NO:21; amino acids Xaa5 through 331 of SEQ ID NO:21; amino acids Xaa5 through

Xaa6 of SEQ ID NO:21, wherein Xaa5 is selected from the group consisting of amino acids 376 through 393 of SEQ ID NO:21 and Xaa6 is selected from the group consisting of amino acids 618 through 629 of SEQ ID NO:21; amino acids 376 through 629 of SEQ ID NO:21; amino acids 393 through 440 of SEQ ID NO:21; amino acids 393 through 618 of SEQ ID NO:21; and amino acids 397 through 611 of SEQ ID NO:21;

- (d) fragments of the amino acid sequences of any of (a)-(c) comprising at least 20 contiguous amino acids;
- (e) fragments of the amino acid sequences of any of (a)-(c) comprising at least 30 contiguous amino acids;
- (f) fragments of the amino acid sequences of any of (a)-(c) having HPR2 polypeptide activity;
- (g) fragments of the amino acid sequences of any of (a)-(c) comprising cytokine receptor domain amino acid sequences;
  - (h) an allelic variant of any of (a)-(c);

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- (i) amino acid sequences comprising at least 20 amino acids and sharing amino acid identity with the amino acid sequences of any of (a)-(h), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99%, and at least 99.5%;
- an amino acid sequence of any of (a)-(i) wherein the polypeptide comprising said (j) amino acid sequence also comprises an amino acid sequence selected from the group consisting of: amino acids 1 through 177 of SEQ ID NO:16; amino acids 216 through 245 of SEQ ID NO:16; SEQ ID NO:17; SEQ ID NO:18; amino acids 349 through 356 of SEQ ID NO:23; amino acids 319 through 565 of SEQ ID NO:21; amino acids Xaa5 through Xaa6 of SEQ ID NO:21, wherein Xaa5 is selected from the group consisting of amino acids 376 through 393 of SEQ ID NO:21 and Xaa6 is selected from the group consisting of amino acids 618 through 629 of SEQ ID NO:21; amino acids 376 through 629 of SEQ ID NO:21; amino acids 393 through 440 of SEQ ID NO:21; amino acids 393 through 618 of SEQ ID NO:21; amino acids 397 through 611 of SEQ ID NO:21; amino acids 381 though 629 of SEQ ID NO:21; a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 comprising at least 20 contiguous amino acids; a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 comprising at least 30 contiguous amino acids; a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 that is at least 25% of the length of the sequence of amino acids 381 though 629 of SEQ ID NO:21; a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 that is at least 50% of the length of the sequence of amino acids 381 though 629 of SEQ ID NO:21; a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 comprising at least one of the following: an HPR2 Box 1 motif, an HPR2 Box 2 motif, and an HPR2 Box 3 motif; and a fragment of the sequence of amino acids 381 though 629 of SEQ ID NO:21 comprising at least one tyrosine residue;

(k) an amino acid sequence of any of (a)-(j) wherein the polypeptide comprising said amino acid sequence does not comprise amino acids 381 through 384 of SEQ ID NO:26;

- (l) an amino acid sequence of (i)-(k), wherein a polypeptide comprising said amino acid sequence of (i)-(k) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (a)-(h); and
- (m) an amino acid sequence of (i)-(l) having HPR2 polypeptide activity.

  Preferably, such polypeptides are isolated HPR2 polypeptides or isolated polypeptides having HPR2 polypeptide activity.

Other aspects of the invention are isolated nucleic acids encoding polypeptides of the invention, with a preferred embodiment being an isolated nucleic acid consisting of, consisting essentially of, or more preferably, comprising a nucleotide sequence selected from the group consisting of:

- (a) SEQ ID NO:19;
- (b) SEQ ID NO:20;
- (c) SEQ ID NO:22;
- (d) SEQ ID NO:24; and
- 20 (d) allelic variants of (a)-(d).

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An additional preferred embodiment of the invention is an isolated nucleic acid consisting of, consisting essentially of, or more preferably, comprising a nucleotide sequence selected from the group consisting of nucleotides 107 through 175 of SEQ ID NO:19, nucleotides 107 through 478 of SEQ ID NO:19, nucleotides 107 through 1060 of SEQ ID NO:19, nucleotides 107 through 1099 of SEQ ID NO:19, nucleotides 176 through 478 of SEQ ID NO:19, nucleotides 176 through 478 of SEQ ID NO:19, nucleotides 176 through 1099 of SEQ ID NO:19, nucleotides 176 through 1099 of SEQ ID NO:19, nucleotides 479 through 1099 of SEQ ID NO:19, nucleotides 479 through 1099 of SEQ ID NO:19, nucleotides 503 through 1033 of SEQ ID NO:19, nucleotides 776 through 1066 of SEQ ID NO:19, nucleotides 776 through 1099 of SEQ ID NO:19, nucleotides 1232 through 1993 of SEQ ID NO:19, nucleotides 1283 through 1426 of SEQ ID NO:19, nucleotides 1283 through 1960 of SEQ ID NO:19, and nucleotides 1295 through 1939 of SEQ ID NO:19.

The invention also provides isolated genomic nucleic acids corresponding to the nucleic acids of the invention.

Another aspect of the invention provides isolated nucleic acids, preferably having a length of at least 15 nucleotides, that hybridize under conditions of moderate stringency to the nucleic acids encoding polypeptides of the invention. In preferred embodiments of the invention, such nucleic acids encode a polypeptide having HPR1 and/or HPR2 polypeptide activity, or comprise a nucleotide sequence that shares nucleotide sequence identity with the nucleotide sequences of the nucleic acids of the invention, wherein the percent nucleotide sequence identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99%. and at least 99.5%.

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Further provided by the invention are expression vectors and recombinant host cells comprising at least one nucleic acid of the invention, and preferred recombinant host cells wherein said nucleic acid is integrated into the host cell genome.

Also provided is a process for producing a polypeptide encoded by the nucleic acids of the invention, comprising culturing a recombinant host cell under conditions promoting expression of said polypeptide, wherein the recombinant host cell comprises at least one nucleic acid of the invention. A preferred process provided by the invention further comprises purifying said polypeptide. In another aspect of the invention, the polypeptide produced by said process is provided.

Further aspects of the invention are isolated antibodies that bind to the polypeptides of the invention, preferably monoclonal antibodies, also preferably humanized antibodies or humanized antibodies, and preferably wherein the antibody inhibits the activity of said polypeptides.

The invention additionally provides a method of designing an inhibitor of the polypeptides of the invention, the method comprising the steps of determining the three-dimensional structure of any such polypeptide, analyzing the three-dimensional structure for the likely binding sites of substrates, synthesizing a molecule that incorporates a predicted reactive site, and determining the polypeptide-inhibiting activity of the molecule.

In a further aspect of the invention, a method is provided for identifying compounds that alter HPR1 and/or HPR2 polypeptide activity comprising

- (a) mixing a test compound with a polypeptide of the invention; and
- (b) determining whether the test compound alters the HPR1 and/or HPR2 polypeptide activity of said polypeptide.

In another aspect of the invention, a method is provided identifying compounds that inhibit the binding activity of HPR1 and/or HPR2 polypeptides comprising

- (a) mixing a test compound with a polypeptide of the invention and a binding partner of said polypeptide; and
- (b) determining whether the test compound inhibits the binding activity of said polypeptide.

In preferred embodiments, the binding partner is a four alpha helix bundle cytokine; more preferably, the binding partner is selected from the group consisting of IL-6, OSM, LIF, CNTF, CLC, IL-12p35, and IL-23p19, and most preferably the binding partners are a soluble hematopoietin receptor such as EBI-3, soluble IL-6R alpha, cytokine-like factor-1 (CLF), IL-12p40, or a soluble form of HPR1 and/or HPR2 in conjunction with a four alpha helix bundle cytokine.

The invention also provides a method for increasing ligand-binding activity, comprising providing at least one compound selected from the group consisting of the polypeptides of the invention and agonists of said polypeptides; with a preferred embodiment of the method further comprising increasing said activity in a patient by administering at least one polypeptide of the invention.

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Further provided by the invention is a method for decreasing ligand-binding activity, comprising providing at least one antagonist of the polypeptides of the invention; with a preferred embodiment of the method further comprising decreasing said activity in a patient by administering at least one antagonist of the polypeptides of the invention, and with a further preferred embodiment wherein the antagonist is an antibody that inhibits the activity of any of said polypeptides.

The invention additionally provides a method for treating a cell proliferation condition comprising administering at least one compound selected from the group consisting of the polypeptides of the invention and agonists of said polypeptides; with a preferred embodiment wherein the cell proliferation condition is selected from the group consisting of pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, and osteoporosis resulting from a lack of bone-forming cells.

The invention additionally provides a method for treating a metabolic condition comprising administering at least one compound selected from the group consisting of the polypeptides of the invention and agonists of said polypeptides; with a preferred embodiment wherein the metabolic condition is obesity.

The invention additionally provides a method for treating a reproductive hormone condition comprising administering at least one compound selected from the group consisting of the polypeptides of the invention and agonists of said polypeptides; with a preferred embodiment wherein the condition is selected from the group consisting of deficient mammary development and infertility.

In other aspects of the invention, a method is provided for treating a cell proliferation condition comprising administering an antagonist of the polypeptide of the invention; with a preferred embodiment wherein the cell proliferation condition is selected from the group consisting of leukemia, tumour metastasis, and osteoporosis resulting from an excess of bone-resorbing cells.

In other aspects of the invention, a method is provided for treating a metabolic condition comprising administering an antagonist of the polypeptide of the invention; with a preferred embodiment wherein the metabolic condition is selected from the group consisting of cachexia, wasting, and AIDS-related weight loss.

In other aspects of the invention, a method is provided for treating cancer conditions stimulated by reproductive hormones comprising administering an antagonist of the polypeptide of the invention; with a preferred embodiment wherein the condition is selected from the group consisting of breast cancer and prolactinoma.

In another embodiment of the invention, methods are provided for using HPR1 and HPR2 polypeptides and antagonists thereof as adjuvants.

A further embodiment of the invention provides a use for the polypeptides of the invention in the preparation of a medicament for treating a cell proliferation condition; with a preferred embodiment wherein the cell proliferation condition is selected from the group consisting of pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, and osteoporosis.

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A further embodiment of the invention provides a use for the polypeptides of the invention in the preparation of a medicament for treating a metabolic condition; with a preferred embodiment wherein the metabolic condition is obesity.

A further embodiment of the invention provides a use for the polypeptides of the invention in the preparation of a medicament for treating a reproductive hormone condition; with a preferred embodiment wherein the condition is selected from the group consisting of deficient mammary development and infertility.

## **DETAILED DESCRIPTION OF THE INVENTION**

#### Similarities of HPR1 and HPR2 Structure to Other Hematopoietin Receptor Family Members

We have identified HPR1 and HPR2, new human hematopoietin receptor polypeptides having structural features characteristic of this polypeptide family; the amino acid sequence of an HPR1 polypeptide is provided in SEQ ID NO:4 and the amino acid sequence of three alternatively spliced forms of HPR2 polypeptide are provided in SEQ ID NOs 21, 23, and 25. We have also identified the murine homologue of human HPR1; the amino acid sequence of Mus musculus HPR1 is presented in SEQ ID NO:12. (The use of "HPR1" without a species designation refers to HPR1 polypeptides generally, for example, human and/or murine, mammalian, or vertebrate HPR1 polypeptides.) Alignments showing the sequence similarities between HPR1, HPR2, and other hematopoietin receptors are presented in Tables 1, 2, and 3 in Example 1 below.

The typical structural elements common to members of the hematopoietin receptor polypeptide family include an extracellular region comprising at least one cytokine receptor domain, and in most members of the family, a cytoplasmic region that in at least a subset of the hematopoietin receptor polypeptides comprises domains involved in intracellular signaling functions. A signal sequence is found at the N-terminus of hematopoietin receptor family polypeptides, and is followed, in N-to-C order, by an immunoglobulin (Ig)-like domain (in some members of the family), a cytokine receptor domain, three copies of a fibronectin repeat (in some members of the family), a transmembrane domain or a glycosyl-phosphatidyl inositol (GPI) linkage to the membrane (except in soluble members of the family, which in most cases are soluble splice variant forms of transmembrane or membrane-linked hematopoietin receptor polypeptides), and a cytoplasmic domain (which is not present in soluble forms). The extracellular domain of hematopoietin receptor polypeptides extends from the N terminus to the transmembrane domain of the protein, and includes the cytokine receptor domain and any Ig-like domains (approximately 100 amino acids in length) or fibronectin repeats (such as fibronectin type III repeats which are approximately 81-83 amino acids in length and are separated by spacer sequences of approximately 10 to 13 amino acids) that may be present in certain of the hematopoietin receptor polypeptides. There are key residues within the cytokine receptor domain, the two or four conserved cysteine residues and the WSXWS motif; substitutions of these residues are likely to be associated with an altered function or lack of that function for the polypeptide. The cytokine receptor domain, which is approximately 200 amino acids in length, can be subdivided into two roughly equal subdomains - an N-terminal 'conserved cysteine' domain and a more C-terminal

5 'WSXWS' domain - separated by a proline-rich 'linker' stretch of four amino acids that allows the two subdomains to form a ligand binding site between them (Bravo and Heath, 2000, *EMBO J.* 19(11): 2399-2411).

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The intracellular domain (also called "cytoplasmic domain") of the hematopoietin receptor polypeptides (in those family members that contain such a domain), extends from the transmembrane domain of the protein to the C terminus, and in the signaling receptor subgroup, includes regions involved in intracellular signal transduction functions. Although the amino acid sequence of the intracellular domain varies considerably between hematopoietin receptor polypeptides, there are a few regions that show some similarity between the members of the family and which have been determined to be involved in binding to members of the signal transduction cascade. "Box 1" is a stretch of 9 to 12 amino acids that begins about 9 amino acids C-terminal to the transmembrane domain, and has within it a conserved Ar-P-X-Al-P-X-P motif, where Ar is an aromatic amino acid (Trp, Phe, or Tyr) and Al is an aliphatic amino acid (Ala, Gly, Val, Leu, or Ile). About 8 amino acids C-terminal to Box 1 there is a conserved aromatic amino acid (usually Trp but also Phe or Tyr), and approximately 15 to 60 amino acids further C-terminal there is a motif of about 11 to 13 amino acids, "Box 2". While Box 1 is present in most of the hematopoietin receptor polypeptides, the Box 2 motif is present in a subset of the hematopoietin receptor family including gp130, GCSFR, LIF-R, the erythropoietin receptor (EPO-R), and several others. Mutations to residues within Box 1 or Box 2, or to the conserved aromatic residue between the Box 1 and Box 2 motifs, have inactivated the ability of the mutated receptor to stimulate cell proliferation upon the addition of ligand. A further conserved domain has been identified in the cytoplasmic domains of signaling cytokine receptors such as gp130, LIF-R, and G-CSFR: "Box 3". The Box 3 motif is about 10 to 15 amino acids located between approximately 70 and 150 amino acids C-terminal of the transmembrane domain, and has a rough match to a (P/T)VXGXGYXXQ consensus sequence. Cytoplasmic regions of these receptors containing Box 3 have been associated with a macrophage differentiation promoting activity (in the case of gp130) and a granulocyte differentiation promoting activity (in the case of G-CSFR) (Soede-Bobok and Touw, 1997, J Mol Med 75: 470-477); however, members of the LIF/IL-6 gp130-sharing family of hematopoietin receptors can also be involved in suppression of differentiation (see Ernst et al., 1999, J Biol Chem 274(14): 9729-9737). Finally, the cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation. Although hematopoietin receptors themselves do not generally have a protein kinase activity, they interact with and are phosphorylated by kinases within the JAK/STAT signal transduction pathways. Mutations in the Box 1 motif abolish the ability of certain of the signaling hematopoietin receptors to bind members of the Janus kinase (JAK) family, particularly JAK2 or JAK1 (Taner et al., 1995, J Biol Chem 270(12): 6523-6530). Hematopoietin receptor-ligand interactions also activate the ERK/MAPK pathway, most likely through the phosphorylation of tyrosine residues in the cytoplasmic domains as the tyrosines at cytoplasmic positions 118 of gp130 (amino acid 759 of SEQ ID NO:8) and 115 of LIF-R (amino acid 974 of SEQ ID NO:6) are present within SHP2 binding sites (Schiemann et al., 1997, J Biol Chem 272(26): 16631-

5 16636). The cytoplasmic tyrosine residues of signaling hematopoietin receptors and the amino acids around them are also important motifs for the recruitment and phosphorylation of signal-transducing STAT polypeptides (Hirano *et al.*, 2000, *Oncogene* 19: 2548-2556).

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Human HPR1 polypeptide has a signal sequence extending from approximately amino acid 20 through amino acid 32 of SEQ ID NO:4, with the mature polypeptide produced by cleavage of this signal sequence predicted to have an amino acid sequence beginning at amino acid 33 of SEQ ID NO:4. Human HPR1 has a cytokine receptor domain extending approximately from amino acid 33 through amino acid 241 of SEQ ID NO:4; three fibronectin repeats from approximately amino acid 242 of SEO ID NO:4 to about amino acid 515 of SEQ ID NO:4; a transmembrane domain that begins approximately between amino acids 526 and 533 of SEQ ID NO:4 and extends to approximately between amino acids 552 and 556 of SEQ ID NO:4 (defining a smaller 'core' transmembrane domain from amino acid 533 to amino acid 552 of SEQ ID NO:4 and an extended transmembrane domain from amino acid 526 to amino acid 556 of SEQ ID NO:4); and a cytoplasmic domain extending from the end of the transmembrane domain (i.e. beginning roughly between amino acids 553 and 557 of SEQ ID NO:4) and extending through the carboxyl terminus of the polypeptide (amino acid 745 of SEQ ID Therefore, human HPR1 polypeptide has an overall structure consistent with other hematopoietin receptor family members. The four conserved cysteine residues within the human HPR1 cytokine receptor domain are located at positions 43, 53, 81, and 94 of SEQ ID NO:4, and the human HPR1 WSXWS motif is located from amino acid 224 through amino acid 228 of SEQ ID NO:4. The human HPR1 N-terminal cytokine receptor subdomain containing four conserved cysteine residues extends approximately from amino acid 33 of SEQ ID NO:4 to amino acid 134 of SEQ ID NO:4; the proline-rich linker is amino acids 135 through 138 of SEQ ID NO:4; and the WSXWS-containing Cterminal cytokine receptor subdomain extends from amino acid 139 to about amino acid 241 of SEQ ID NO:4. In human HPR1, as in several members of the hematopoietin receptor family, the cytokine receptor domain is followed by three fibronectin type III repeats; these repeats are located within the human HPR1 amino acid sequence of SEQ ID NO:4 at the following approximate locations: amino acids 242 to 244 through 324 to 326, amino acids 336 to 337 through 419 to 422, and amino acids 430 to 433 through 514 to 515. Within its intracellular domain, human HPR1 polypeptide contains a good match to the Box 1 conserved motif from amino acid 563 through amino acid 573 of SEQ ID NO:4, a conserved downstream Trp residue (amino acid 581 of SEQ ID NO:4), and a Box 2 motif from amino acid 631 to amino acid 641 of SEQ ID NO:4. The cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation; in human HPR1, such tyrosines are located at positions 652, 683, and 721 of SEQ ID NO:4. Human HPR1 contains several instances of an Asp-containing motif within its cytoplasmic region. In the area overlapping the Box 2 location, human HPR1 has repeated amino acid sequences as shown in the following table; these sequences form a consensus sequence of DKL(N/V)(T/Al), where Al is an aliphatic residue as described above. Other signaling hematopoietin receptors such as murine HPR1 (at

5 amino acids 600 through 604 of SEQ ID NO:12) and gp130 also contain at least one similar Aspcontaining sequence in the region around and following the Box 2 location.

Repeat Sequence	Location in SEQ ID NO:4
DKLNL	amino acids 588 through 592
DSVNT	amino acids 597 through 601
DRILK	amino acids 603 through 607
DKLVI	amino acids 614 through 618
DKLVV	amino acids 619 through 623
DEART	amino acids 635 through 639

Variants, presumably splice variants, of human HPR1 are described in WO 00/75314: a 252-amino-acid form ("NR10.2"), a 652-amino-acid form ("NR10.1"), and a 662-amino-acid form ("NR10.3"). The 252-amino-acid form of HPR1 (SEQ ID NO:13) is identical to SEQ ID NO:4 through amino acid 238, and then has a divergent amino acid sequence from amino acid 239 through 252 of SEQ ID NO:13. This 252-amino-acid form of human HPR1 therefore does not contain the fibronectin type III repeats found in the full-length 745-amino-acid HPR1 of SEQ ID NO:4, or the transmembrane domain or the intracellular region of the SEQ ID NO:4 polypeptide. The 652-amino-acid form of HPR1 (SEQ ID NO:14) is identical to SEQ ID NO:4 through amino acid 642, and then has a divergent amino acid sequence from amino acid 643 through 652 of SEQ ID NO:14.; and the 662-amino-acid form of HPR1 (SEQ ID NO:15) is identical to SEQ ID NO:4 through amino acid 651, and then has a divergent amino acid sequence from amino acid 652 through 662 of SEQ ID NO:15. The 652- and 662-amino-acid forms of human HPR1 therefore do not contain the tyrosine residues at positions 652, 683, and 721 of the intracellular region of the SEQ ID NO:4 polypeptide which are potential substrates for phosphorylation by kinases, such as those of the ERK/MAPK signaling pathways.

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The Mus musculus HPR1 amino acid sequence of SEQ ID NO:12 has a signal sequence beginning approximately between amino acid 13 and amino acid 16 of SEQ ID NO:12 and extending approximately through amino acid 28 of SEQ ID NO:12, with the mature polypeptide produced by cleavage of this signal sequence predicted to have an amino acid sequence beginning at amino acid 29 of SEQ ID NO:12. Murine HPR1 has a cytokine receptor domain extending approximately from amino acid 29 through amino acid 224 of SEQ ID NO:12; three fibronectin repeats from approximately amino acid 225 of SEO ID NO:12 to about amino acid 499 of SEO ID NO:12; a transmembrane domain that begins approximately between amino acids 510 and 517 of SEQ ID NO:12 and extends to approximately between amino acids 532 and 533 of SEQ ID NO:12 (defining a smaller 'core' transmembrane domain from amino acid 517 to amino acid 532 of SEQ ID NO:12 and an extended transmembrane domain from amino acid 510 to amino acid 533 of SEQ ID NO:12); and a cytoplasmic domain extending from the end of the transmembrane domain (i.e. beginning roughly between amino acids 533 and 534 of SEQ ID NO:12) and extending through the carboxyl terminus of the polypeptide (amino acid 726 of SEQ ID NO:12). Therefore, murine HPR1 polypeptide has an overall structure consistent with other hematopoietin receptor family members. There are two conserved cysteine residues within the murine HPR1 cytokine receptor domain located at positions 39 and 49of SEQ ID NO:12, and there are two additional cysteines in this region (although at non-conserved positions) at

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amino acids 90 and 97 of SEQ ID NO:12. The murine HPR1 WSXWS motif is located from amino acid 207 through amino acid 211 of SEQ ID NO:12. The murine HPR1 N-terminal cytokine receptor subdomain containing two conserved cysteine residues (and two additional cysteine residues) extends approximately from amino acid 29 of SEQ ID NO:12 to amino acid 124 of SEQ ID NO:12; the prolinerich linker is amino acids 125 through 128 of SEQ ID NO:12; and the WSXWS-containing C-terminal cytokine receptor subdomain extends from amino acid 129 to about amino acid 224 of SEQ ID NO:12. In murine HPR1, as in several members of the hematopoietin receptor family, the cytokine receptor domain is followed by three fibronectin type III repeats; these repeats are located within the murine HPR1 amino acid sequence of SEQ ID NO:12 at the following approximate locations: amino acids 225 to 227 through 307 to 309, amino acids 319 to 320 through 403 to 406, and amino acids 413 to 417 through 498 to 499. Within its intracellular domain, murine HPR1 polypeptide contains a good match to the Box 1 conserved motif from amino acid 547 through amino acid 557 of SEQ ID NO:12, a conserved downstream Trp residue (amino acid 565 of SEQ ID NO:12), and a Box 2 motif from amino acid 612 through amino acid 622 of SEQ ID NO:12. The cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation; in murine HPR1, such tyrosines are located at positions 633, 674, and 701 of SEQ ID NO:12.

Human HPR2 polypeptide has a signal sequence extending from approximately amino acid 11 through amino acid 23 of SEQ ID NO:21, with the mature polypeptide produced by cleavage of this signal sequence predicted to have an amino acid sequence beginning at amino acid 24 of SEQ ID NO:21. The membrane-spanning (629 amino acids) form of HPR2 has an N-terminal Ig-like domain extending approximately from amino acid 24 through amino acid 124 of SEQ ID NO:21, a cytokine receptor domain extending approximately from amino acid 125 through an amino acid from 320 to 331 of SEQ ID NO:21; a transmembrane domain that begins approximately at amino acid 356 of SEQ ID NO:21 and extends to approximately amino acid 375 of SEQ ID NO:21; and a cytoplasmic domain extending from the end of the transmembrane domain (i.e. beginning approximately at amino acid 376 of SEQ ID NO:21) and extending through the carboxyl terminus of the polypeptide (amino acid 629 of SEQ ID NO:21). Therefore, HPR2 polypeptide has an overall structure consistent with other hematopoietin receptor family members. The N-terminal Ig-like domain contains six cysteine residues at positions 30, 52, 59, 101, 105, and 115 of SEQ ID NO:21, the most conserved of which appear to be the two cysteines at positions 52 and 101; the cysteines at positions 30, 115 (and to a lesser extent, at 105) also align with cysteines at similar positions in Ig or Ig-like domains. The HPR2 Ig-like domain appears to have the greatest degree of sequence similarity with members of the LIR (leukocyte Ig-like receptor) polypeptide family, particularly LIR-3 and LIR-4. The two conserved cysteine residues within the human HPR2 cytokine receptor domain are located at amino acid positions 133 and 144 of SEQ ID NO:21, and the HPR2 version of the WSXWS motif, which has a glutamine residue at the second position of the motif rather than a serine residue, is located from amino acid 304 through amino acid 308 of SEQ ID NO:21. The HPR2 N-terminal cytokine receptor subdomain containing the two

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conserved cysteine residues extends approximately from amino acid 125 of SEQ ID NO:21 to amino acid 219 of SEQ ID NO:21; the proline-rich linker (in this case, proline- and alanine-rich) is amino acids 220 through 223 of SEQ ID NO:21; and the 'WQXWS'-containing C-terminal cytokine receptor subdomain extends from amino acid 224 through an amino acid from 320 to 331 of SEQ ID NO:21. HPR2 does not contain the fibronectin type III repeats found in human and murine HPR1. Within its intracellular domain, the membrane-spanning (629 amino acids) form of HPR2 contains a good match to the Box 1 conserved motif from amino acid 393 through amino acid 403 of SEQ ID NO:21, does not contain a Trp residue between Box 1 and Box2, and has a Box 2 motif from amino acid 430 to amino acid 440 of SEQ ID NO:21. There are also two matches to the Box 3 motif in this membrane-spanning HPR2 polypeptide, at amino acids 478 through 491 and at amino acids 605 through 618 of SEQ ID NO:21. The cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation; in human HPR2, such tyrosines are located at amino acid positions 397 (within the Box 1 motif), 429 (immediately N-terminal to the Box 2 motif), 450, 463, and 476 (just N-terminal of the most N-terminal Box 3 motif), and amino acids 484 and 611 (each of these last two amino acids is within a Box 3 motif) of SEQ ID NO:21. In several respects, the membrane-spanning form of HPR2 shows similarity to the LIF-R hematopoietin receptor: both of these molecules have an Ig-like domain that is followed by a cytokine receptor domain having two (as compared to four) conserved cysteines; and both have Box 1, Box 2, and Box 3 motifs in their intracellular domains, and do not have a tryptophan residue between Box 1 and Box 2.

The HPR2-ex9 polypeptide of SEQ ID NO:23 (356 amino acids), created by alternative splicing which removes exon 9 of the HPR2 coding sequence (see Example 1 below), is identical to the HPR2 629-amino-acid form from amino acid 1 through amino acid 348, but then diverges in sequence for the eight amino acids from amino acid 349 to the C terminus at amino acid 356. The HPR2-ex9 form does not contain a transmembrane region, and is expected to be a secreted form of HPR2 containing the HPR2 extracellular Ig-like and cytokine receptor domains. The HPR2-ex8-ex9 polypeptide of SEQ ID NO:25 (565 amino acids), created by alternative splicing which removes exons 8 and 9 of the HPR2 coding sequence (see Example 1 below), is identical to the HPR2 629-amino-acid form from amino acid 1 through amino acid 318, is missing the next 64 amino acids which include the transmembrane domain, but then shows identity between amino acid 319 through amino acid 565 of SEQ ID NO:25 and the C-terminal region of the 629-amino-acid form of HPR2. The HPR2-ex8-ex9 form is also expected to be a secreted form of HPR2 containing not only the HPR2 extracellular Ig-like and cytokine receptor domains, but also the C-terminal portion of the HPR2 protein which includes the Box 1, Box 2, and Box 3 motifs. A variant, presumably a splice variant, of human HPR2 is described in WO 00/73451: a 384-amino-acid form ("DCRS2"). This 384-amino-acid form of HPR2 (SEQ ID NO:26) is identical to SEQ ID NO:21 through amino acid 380, and then has a divergent amino acid sequence from amino acid 381 through 384 of SEQ ID NO:26. This 384-amino-acid form of human HPR2 therefore does not contain the intracellular region of the SEQ ID NO:21 HPR2 polypeptide,

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which contains the Box1, 2, and 3 motifs and intracellular tyrosine residues that are involved in the signaling (or signal transduction) function of the SEQ ID NO:21 HPR2 polypeptide.

The Mus musculus HPR2 amino acid sequence of SEQ ID NO:27 has a signal sequence beginning approximately between amino acid 8 and amino acid 11 and extending through amino acid 23 of SEQ ID NO:27, with the mature polypeptide produced by cleavage of this signal sequence predicted to have an amino acid sequence beginning at amino acid 24 of SEQ ID NO:27. Mus musculus HPR2, like the membrane-spanning form of human HPR2, has an N-terminal Ig-like domain extending approximately from amino acid 24 through amino acid 124 of SEQ ID NO:27, a cytokine receptor domain extending approximately from amino acid 125 through an amino acid from 341 to 350 of SEQ ID NO:27; a transmembrane domain that begins approximately between amino acid 373 and amino acid 380 of SEO ID NO:27 and extends through approximately between amino acid 394 and amino acid 395 of SEQ ID NO:27 (defining a smaller 'core' transmembrane domain from amino acid 380 to amino acid 394 of SEQ ID NO:27 and an extended transmembrane domain from amino acid 373 to amino acid 395 of SEQ ID NO:27); and a cytoplasmic domain extending from the end of the transmembrane domain (i.e. beginning approximately at amino acid 395 or at amino acid 396 of SEQ ID NO:27) and extending through the carboxyl terminus of the polypeptide (amino acid 644 of SEQ ID Therefore, murine HPR2 polypeptide has an overall structure consistent with other hematopoietin receptor family members. The N-terminal Ig-like domain contains six cysteine residues at positions 30, 52, 59, 101, 105, and 115 of SEQ ID NO:27, the most conserved of which appear to be the two cysteines at positions 52 and 101; the cysteines at positions 30, 115 (and to a lesser extent, at 105) also align with cysteines at similar positions in Ig or Ig-like domains. As with human HPR2, the murine HPR2 Ig-like domain appears to have the greatest degree of sequence similarity with members of the LIR (leukocyte Ig-like receptor) polypeptide family. The two conserved cysteine residues within the human HPR2 cytokine receptor domain are located at amino acid positions 133 and 144 of SEQ ID NO:27, and the murine HPR2 version of the "WSXWS" motif, which like human HPR2 has a glutamine residue at the second position of the motif rather than a serine residue, is located from amino acid 324 through amino acid 328 of SEQ ID NO:27. The murine HPR2 polypeptide contains an insert of 20 amino acids relative to the human HPR2 polypeptide; this insert region extends from amino acid 297 through amino acid 316 of SEQ ID NO:27, and is a perfect repeat of amino acids 317 through 336 of SEQ ID NO:27. Therefore, in the SEQ ID NO:27 form of murine HPR2, there is a second WQXWS motif at amino acids 304 through 308 of SEQ ID NO:27. The murine HPR2 N-terminal cytokine receptor subdomain containing the two conserved cysteine residues extends approximately from amino acid 125 of SEQ ID NO:27 to amino acid 219 of SEQ ID NO:27; the proline-rich linker (in this case, proline- and alanine-rich) is amino acids 220 through 223 of SEO ID NO:27; and the C-terminal cytokine receptor subdomain containing the two repeats of the WQXWS motif extends from amino acid 224 through an amino acid from 340 to 350 of SEQ ID NO:27. Murine HPR2 does not contain the fibronectin type III repeats found in human and murine HPR1. Within its intracellular domain, this membrane-spanning form of murine HPR2 contains a good match to the Box 1 conserved motif from

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amino acid 412 through amino acid 422 of SEQ ID NO:27, does not contain a Trp residue between Box 1 and Box2, and has a Box 2 motif from amino acid 449 to amino acid 459 of SEQ ID NO:27. There are also two matches to the Box 3 motif in this murine membrane-spanning HPR2 polypeptide, at amino acids 498 through 511 and at amino acids 620 through 633 of SEQ ID NO:27. The cytoplasmic domains of signaling hematopoietin receptor polypeptides contain several tyrosine residues that are potential sites for phosphorylation; in murine HPR2, such tyrosines are located at amino acid positions 416 (within the Box 1 motif), 448 (immediately N-terminal to the Box 2 motif), 469, and 496 (just N-terminal of the most N-terminal Box 3 motif), and amino acids 504 and 626 (each of these last two amino acids is within a Box 3 motif) of SEQ ID NO:27. There is an additional intracellular tyrosine located at position 542 of SEQ ID NO:27. As with the membrane-spanning form of human HPR2, murine HPR2 shows similarity to the LIF-R hematopoietin receptor.

Each of the HPR1 and the HPR2 groups of related polypeptides therefore contains a distinct subset of the several features characteristic of at least some members of the hematopoietin receptor family. The skilled artisan will recognize that the boundaries of the regions of the HPR1 and HPR2 polypeptides described above are approximate and that the precise boundaries of such domains, as for example the boundaries of the transmembrane region (which can be predicted by using computer programs available for that purpose), can also differ from member to member within the hematopoietin receptor polypeptide family.

The hematopoietin receptor polypeptide family is highly to moderately conserved between species, with the family members within a particular species exhibiting some sequence conservation, particularly with respect to the conserved domains and residues described above. Subfamilies of the hematopoietin receptor polypeptide family can be defined on the basis of structure, for example the Iglike domain containing members, or the fibronectin repeat containing members. It is also possible to group hematopoietin receptor polypeptides according to the length of the cytoplasmic domain, with those receptors having a longer cytoplasmic domain being more likely to be signaling receptors. Subgroups of the hematopoietin receptor family can also be defined on the basis of a shared common signaling receptor present in several different combinations of heteromeric receptors. For example, the gp130 signaling receptor is found in separate complexes with LIF-R, IL-6R alpha or a soluble form of IL-6R alpha, and CNTFR alpha; monomeric forms or multimeric combinations of these receptor components bind to IL-6, OSM, LIF, and/or CNTF; thus a "gp130-sharing group" subfamily would include these hematopoietin receptor polypeptides and be associated with this group of cytokines. Another group of hematopoietin receptors are those which associate with a ligand comprising at least two soluble polypeptides. For example, the IL-12 receptor associates with the combination of the p40 polypeptide, similar in structure to soluble forms of hematopoietin receptors such as soluble IL-6R alpha, and the four alpha helix bundle p35 polypeptide. The IL-12 p40 subunit can also associate with another four alpha helix bundle cytokine called p19; when p40 binds p19 the resulting combination has been named "IL-23" and has been shown to bind to the IL-12R beta 1 receptor subunit, but not the signaling IL-12R beta 2 receptor subunit (Oppmann et al., 2000, Immunity 13: 715-725). Thus the

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p40-p19 complex is likely to bind a different IL-12RB2-like signaling receptor subunit, such as HPR2, HPR1, GCSFR, or gp130. As another example, CNTFR alpha, gp130, and LIFR can each associate with a combination of the soluble receptor cytokine-like factor-1 (CLF-1) and cardiotrophin-like cytokine (CLC), with CLF-1 and CLC analogous to p40 and p35, respectively (Elson et al., 2000, Nat Neurosci 3(9): 867-872). The cytokine receptor domains of HPR1 and HPR2 are similar in sequence to those of gp130, IL-6R beta, IL-12RB2, GCSFR, LIFR, leptin receptor, prolactin receptor, and other members of the hematopoietin receptor family, with HPR1 showing the greatest degree of similarity to gp130 and IL-6R beta, and HPR2 showing the greatest degree of similarity to gp130 and IL-12RB2. Because HPR1 and HPR2 each have a substantial cytoplasmic domain and are most similar in sequence to gp130, HPR1 and HPR2 are likely to be new signaling members of the "gp130-sharing" subfamily of hematopoietin receptors; however, HPR2 may also share attributes of the IL-12RB2 receptor subunit, such as involvement in modulation of the balance between Th1 and Th2 immune responses. Expression of HPR1 and HPR2 has been detected by PCR amplification from tissue-specific cDNA libraries in several cell types including COS-1 cells, 293MSR cells, the B cell lines CB23 and MP-1, the B cell lymphoma lines Daudi, and Raji, the T cell leukemia line HSB2, and the promonocytic leukemia line U937. HPR2 mRNA expression appears to be more prevalent than HPR1 expression in the B cell derived lines, while HPR1 mRNA expression appears to be more prevalent than HPR2 expression in the T cell derived and monocyte lines. EBI-3 is a p40-like soluble hematopoietin receptor polypeptide; FACS analysis has shown that EBI-3-Fc fusion polypeptides bind to cells expressing HPR1 and HPR2 such as COS-1 cells, 293MSR cells, and CB23 and MP-1 cells, indicating that EBI-3 is a potential binding partner of HPR1 and HPR2, most likely in conjunction with a four alpha helix bundle cytokine such as IL-6, OSM, LIF, CNTF, CLC, IL-12p35, or IL-23p19.

### Biological Activities and Functions of HPR1 and HPR2 Polypeptides

PCR amplification from tissue-specific cDNA libraries was performed to detect HPR1 or HPR2 cDNA sequences. The results of these experiments show that HPR1 transcripts are expressed in a wide variety of fetal and adult human cells, including testis, lung, placenta, pancreas, prostate, peripheral blood cells, thymus, stomach, and skin cells; as well as in various cell lines including U937 cells, the leukemia cell line HSB2, LX-1/GI-117 lung carcinoma cells, GI-112 colon adenocarcinoma cells, the B cell lines MP-1 and CB23, COS-1 cells, and 293MSR cells. HPR2 transcripts are present in a similarly diverse group of adult and fetal human cell types, including placenta, lung, kidney, pancreas, prostate, testis, colon, LX-1/GI-117 lung carcinoma cells, tonsil/CX-1 cells, lymph node, GI-112 colon adenocarcinoma cells, heart, brain, spleen, thymus, ovary, small intestine, fetal brain, fetal lung/heart, fetal spleen, fetal thymus, esophagus, stomach, and skin; and in various cell lines such as the B cell lines MP-1 and CB23, Daudi cells, Raji cells, HSB2 cells, COS-1 cells, and 293MSR cells.

Typical biological activities or functions associated with HPR1 and HPR2 polypeptides are ligand-binding activity, intracellular signaling activity, cell proliferation stimulatory activity, cell proliferation inhibitory activity, cell differentiation stimulatory activity, and cell differentiation

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inhibitory activity. HPR1 and HPR2 polypeptides having ligand-binding activity bind to cytokine or growth factor ligand molecules of the four alpha helix bundle family of cytokines, and in particular are likely to bind cytokines such as IL-6, OSM, LIF, CNTF, CLC, IL-12p35, and IL-23p19, and/or soluble hematopoietin receptors such as EBI-3, soluble IL-6R alpha, cytokine-like factor-1 (CLF), IL-12p40, or a soluble form of HPR1 and/or HPR2. This ligand-binding activity is associated with the extracellular cytokine receptor domain of HPR1 polypeptides. Thus, for uses requiring ligand-binding activity, preferred HPR1 and HPR2 polypeptides include those having at least one cytokine receptor domain and exhibiting ligand-binding activity. Preferred HPR1 and HPR2 polypeptides further include oligomers or fusion polypeptides comprising at least one cytokine receptor portion of one or more HPR1 and/or HPR2 polypeptides, and fragments of any of these polypeptides that have ligand-binding activity. The ligand-binding activity of HPR1 and HPR2 polypeptides may be determined, for example, by any standard assay to measure binding of labeled ligand or by a competitive binding assay, all of which are described more extensively below. HPR1 and HPR2 polypeptides having intracellular signaling activity bind ligand molecules when in association with other receptor polypeptides to form a homo- or heteromeric complex, with ligand binding initiating a signaling cascade. The intracellular signaling activity is associated with the cytoplasmic domain of certain HPR1 and HPR2 polypeptides. Thus, for uses requiring intracellular signaling activity, preferred HPR1 and HPR2 polypeptides include those having the cytoplasmic domain, and in particular having certain conserved domains (such as the Box 1 motif, the Trp residue at position 581 of SEQ ID NO:4, the Box 2 motif, the Asp-containing motifs between amino acids 588 and 639 of SEQ ID NO:4, or the Box 3 motif) and conserved cytoplasmic tyrosine residues, and exhibiting intracellular signaling biological activity. Preferred HPR1 and HPR2 polypeptides further include oligomers or fusion polypeptides comprising at least one cytoplasmic portion of one or more HPR1 and/or HPR2 polypeptides, and fragments of any of these polypeptides that have intracellular signaling activity. The intracellular signaling activity of HPR1 and HPR2 polypeptides may be determined, for example, through assays to detect phosphorylation of the HPR1 polypeptide, the HPR2 polypeptide, or downstream polypeptides in signaling cascades such as the JAK/STAT or ERK/MAPK pathways, or in assays that measure biological activities related to the signal transmission, such as stimulation or suppression of cell proliferation, differentiation, or activation. One example of an assay to measure cytokine-binding and cell-proliferation activity involves expressing a polypeptide of the invention in Ba/F3 cells, exposing the polypeptide-expressing cells to radioactively labeled cytokine, and measuring specific cytokine binding to cells and uptake of 3H-thymidine by cells in response to cytokine, as described in Presky et al., 1996, Proc Natl Acad Sci USA 93: 14002-14007. Further examples of such assays are described herein and in Ernst et al., 1999, J Biol Chem 274(14): 9729-9737. Soluble forms of hematopoietin receptors comprising one or more extracellular domains of the hematopoietin receptor, such as soluble forms of HPR1 and HPR2, may also be used in assays to measure their effect on cell growth, proliferation, differentiation, or activation; in such assays the cells are contacted with the soluble form of the receptor and their growth,

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proliferation, differentiation, or activation is measured, for example by measuring the incorporation of radioactive thymidine or by microscopic examination of treated and untreated cells.

The terms "HPR1 polypeptide activity" and "HPR2 polypeptide activity," as used herein, include any one or more of the following: ligand-binding activity and intracellular signaling activity (which includes effects on cell growth, proliferation, differentiation, or activation), as well as the *ex vivo* and *in vivo* activities of HPR1 and HPR2 polypeptides. The degree to which HPR1 and HPR2 polypeptides and fragments and other derivatives of these polypeptides exhibit these activities can be determined by standard assay methods as disclosed herein; those of skill in the art will appreciate that other, similar types of assays can be used to measure HPR1 and HPR2 biological activities.

Another aspect of the biological activity of HPR1 and HPR2 polypeptides is the ability of members of these polypeptide families to bind particular binding partners such as cytokines, other hematopoietin receptor polypeptides, and intracellular signaling polypeptides, with the cytokine receptor domain binding to cytokines and the intracellular signaling domain binding to intracellular signaling polypeptides such as members of the JAK and SHP polypeptide families. The term "binding partner," as used herein, includes ligands, receptors, substrates, antibodies, other hematopoietin receptor polypeptides, the same HPR1 or HPR2 polypeptide (in the case of homotypic interactions), and any other molecule that interacts with an HPR1 or an HPR2 polypeptide through contact or proximity between particular portions of the binding partner and the HPR1 or HPR2 polypeptide. Because the cytokine receptor domains of HPR1 and HPR2 polypeptides bind to cytokines, an HPR1 or HPR2 cytokine receptor domain when expressed as a separate fragment from the rest of an HPR1 or HPR2 polypeptide, or as a soluble polypeptide, fused for example to an immunoglobulin Fc domain, is expected to disrupt the binding of endogenous HPR1 and/or HPR2 polypeptides to their binding partners. By binding to one or more binding partners, the separate cytokine receptor domain polypeptide likely prevents binding by the native HPR1 and/or HPR2 polypeptide(s), and so acts in a dominant negative fashion to inhibit the biological activities mediated via binding of HPR1 and/or HPR2 polypeptides to cytokines. Assays for evaluating the biological activities and partner-binding properties of HPR1 and HPR2 polypeptides are described further herein.

HPR1 and HPR2 polypeptides are involved in cell proliferation, differentiation, or activation diseases or conditions, that share as a common feature ligand-binding activity in their etiology. More specifically, the following cell proliferation conditions are those that are known or are likely to involve the biological activities of HPR1 and/or HPR2 polypeptides: pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, osteoporosis resulting from a lack of bone-forming cells, leukemia, tumour metastasis, and osteoporosis resulting from an excess of bone-resorbing cells. In addition, the following metabolic conditions involving hematopoietin receptor ligands such as leptin are those that are known or are likely to involve the biological activities of HPR1 and/or HPR2 polypeptides: obesity, cachexia, wasting, and AIDS-related weight loss. Also, the following prolactin-related conditions are those that are known or are likely to involve the biological activities of HPR1 and/or HPR2 polypeptides: deficient mammary development, infertility, breast cancer, and

prolactinoma. Blocking or inhibiting the interactions between members of the HPR1 and HPR2 polypeptide families and their substrates, ligands, receptors, binding partners, and or other interacting polypeptides is an aspect of the invention and provides methods for treating or ameliorating these diseases and conditions through the use of inhibitors of HPR1 and/or HPR2 polypeptide activity. Examples of such inhibitors or antagonists are described in more detail below. For certain conditions involving too little HPR1 or HPR2 polypeptide activity, methods of treating or ameliorating these conditions comprise increasing the amount or activity of HPR1 or HPR2 polypeptides by providing isolated HPR1 or HPR2 polypeptides or active fragments or fusion polypeptides thereof, or by providing compounds (agonists) that activate endogenous or exogenous HPR1 or HPR2 polypeptides.

#### **HPR1** and **HPR2** Polypeptides

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An HPR1 polypeptide is a polypeptide that shares a sufficient degree of amino acid identity or similarity to the human HPR1 polypeptide of SEQ ID NO:4 or the murine HPR1 polypeptide of SEQ ID NO:12 to (A) be identified by those of skill in the art as a polypeptide likely to share particular structural domains and/or (B) have biological activities in common with the HPR1 polypeptides of SEQ ID NO:4 and SEQ ID NO:12 and/or (C) bind to antibodies that also specifically bind to other HPR1 polypeptides. An HPR2 polypeptide is a polypeptide that shares a sufficient degree of amino acid identity or similarity to the HPR2 polypeptides of SEQ ID NOs 21, 23, 25, and 27 to (A) be identified by those of skill in the art as a polypeptide likely to share particular structural domains and/or (B) have biological activities in common with the HPR2 polypeptides of SEQ ID NOs 21, 23, 25, and 27 and/or (C) bind to antibodies that also specifically bind to other HPR2 polypeptides. HPR1 and HPR2 polypeptides can be isolated from naturally occurring sources, or have the same structure as naturally occurring HPR1 or HPR2 polypeptides, or can be produced to have structures that differ from naturally occurring HPR1 or HPR2 polypeptides. Polypeptides derived from any HPR1 or HPR2 polypeptide by any type of alteration (for example, but not limited to, insertions, deletions, or substitutions of amino acids; changes in the state of glycosylation of the polypeptide; refolding or isomerization to change its three-dimensional structure or self-association state; and changes to its association with other polypeptides or molecules) are also HPR1 or HPR2 polypeptides, respectively. Therefore, the polypeptides provided by the invention include polypeptides characterized by amino acid sequences similar to those of the HPR1 and HPR2 polypeptides described herein, but into which modifications are naturally provided or deliberately engineered. A polypeptide that shares biological activities in common with members of the HPR1 and/or HPR2 polypeptide family is a polypeptide having HPR1 and/or HPR2 polypeptide activity. Examples of biological activities exhibited by HPR1 and/or HPR2 polypeptides include, without limitation, ligand-binding activity and intracellular signaling.

The present invention provides both full-length and mature forms of HPR1 and HPR2 polypeptides. Full-length polypeptides are those having the complete primary amino acid sequence of the polypeptide as initially translated. The amino acid sequences of full-length polypeptides can be obtained, for example, by translation of the complete open reading frame ("ORF") of a cDNA

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molecule. Several full-length polypeptides can be encoded by a single genetic locus if multiple mRNA forms are produced from that locus by alternative splicing or by the use of multiple translation initiation sites. The "mature form" of a polypeptide refers to a polypeptide that has undergone posttranslational processing steps such as cleavage of the signal sequence or proteolytic cleavage to remove a prodomain. Multiple mature forms of a particular full-length polypeptide may be produced, for example by cleavage of the signal sequence at multiple sites, or by differential regulation of proteases that cleave the polypeptide. The mature form(s) of such polypeptide can be obtained by expression, in a suitable mammalian cell or other host cell, of a nucleic acid molecule that encodes the full-length polypeptide. The sequence of the mature form of the polypeptide may also be determinable from the amino acid sequence of the full-length form, through identification of signal sequences or protease cleavage sites. The HPR1 and HPR2 polypeptides of the invention also include those that result from post-transcriptional or post-translational processing events such as alternate mRNA processing which can yield alternative splice forms of HPR1 or HPR2 such as a truncated but biologically active polypeptide or, for example, a naturally occurring soluble form of the polypeptide. Also encompassed within the invention are variations attributable to proteolysis such as differences in the N- or C-termini upon expression in different types of host cells, due to proteolytic removal of one or more terminal amino acids from the polypeptide (generally from 1-5 terminal amino acids).

The invention further includes HPR1 and HPR2 polypeptides with or without associated native-pattern glycosylation. Polypeptides expressed in yeast or mammalian expression systems (e.g., COS-1 or CHO cells) can be similar to or significantly different from a native polypeptide in molecular weight and glycosylation pattern, depending upon the choice of expression system. Expression of polypeptides of the invention in bacterial expression systems, such as E. coli, provides non-glycosylated molecules. Further, a given preparation can include multiple differentially glycosylated species of the polypeptide. Glycosyl groups can be removed through conventional methods, in particular those utilizing glycopeptidase. In general, glycosylated polypeptides of the invention can be incubated with a molar excess of glycopeptidase (Boehringer Mannheim).

Species homologues of HPR1 and HPR2 polypeptides and of nucleic acids encoding them are also provided by the present invention. As used herein, a "species homologue" is a polypeptide or nucleic acid with a different species of origin from that of a given polypeptide or nucleic acid, but with significant sequence similarity to the given polypeptide or nucleic acid, as determined by those of skill in the art. Species homologues can be isolated and identified by making suitable probes or primers from polynucleotides encoding the amino acid sequences provided herein and screening a suitable nucleic acid source from the desired species. The invention also encompasses allelic variants of HPR1 and HPR2 polypeptides and nucleic acids encoding them; that is, naturally-occurring alternative forms of such polypeptides and nucleic acids in which differences in amino acid or nucleotide sequence are attributable to genetic polymorphism (allelic variation among individuals within a population).

Fragments of the HPR1 and HPR2 polypeptides of the present invention are encompassed by the present invention and can be in linear form or cyclized using known methods, for example, as

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described in Saragovi, et al., Bio/Technology 10, 773-778 (1992) and in McDowell, et al., J. Amer. Chem. Soc. 114 9245-9253 (1992). Polypeptides and polypeptide fragments of the present invention, and nucleic acids encoding them, include polypeptides and nucleic acids with amino acid or nucleotide sequence lengths that are at least 25% (more preferably at least 50%, or at least 60%, or at least 70%, and most preferably at least 80%) of the length of an HPR1 polypeptide or of an HPR2 polypeptide, and have at least 60% sequence identity (more preferably at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, or at least 99%, and most preferably at least 99.5%) with that HPR1 or HPR2 polypeptide or encoding nucleic acid, where sequence identity is determined by comparing the amino acid sequences of the polypeptides when aligned so as to maximize overlap and identity while minimizing sequence gaps. Also included in the present invention are polypeptides and polypeptide fragments, and nucleic acids encoding them, that contain or encode a segment preferably comprising at least 8, or at least 10, or preferably at least 15, or more preferably at least 20, or still more preferably at least 30, or most preferably at least 40 contiguous amino acids. Such polypeptides and polypeptide fragments may also contain a segment that shares at least 70% sequence identity (more preferably at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, or at least 99%, and most preferably at least 99.5%) with any such segment of any of the HPR1 or HPR2 polypeptides, where sequence identity is determined by comparing the amino acid sequences of the polypeptides when aligned so as to maximize overlap and identity while minimizing sequence gaps. The percent identity can be determined by visual inspection and mathematical calculation. Preferably, the comparison is done using a computer program. An exemplary, preferred computer program is the Genetics Computer Group (GCG; Madison, WI) Wisconsin package version 10.0 program, 'GAP.' The preferred default parameters for the 'GAP' program includes: (1) The GCG implementation of comparison matrices for nucleotides and amino acids; such as a unary comparison matrix (containing a value of 1 for identities and 0 for non-identities) for nucleotides, and the weighted comparison matrix of Gribskov and Burgess, Nucl. Acids Res. 14:6745, 1986, as described by Schwartz and Dayhoff, eds., Atlas of Polypeptide Sequence and Structure, National Biomedical Research Foundation, pp. 353-358, 1979; (2) a penalty of 30 for each gap and an additional penalty of 1 for each symbol in each gap for amino acid sequences, or penalty of 50 for each gap and an additional penalty of 3 for each symbol in each gap for nucleotide sequences; (3) no penalty for end gaps; and (4) no maximum penalty for long gaps. Another program useful for determining percent identify is the BESTFIT program, also available from the University of Wisconsin as part of the GCG computer package. Default parameters for using the BESTFIT program are the same as those described above for using the GAP program. Other programs used by those skilled in the art of sequence comparison can also be used, such as, for example, the UW-BLAST 2.0 algorithm or the BLASTN program version 2.0.9, available for use via the National Library of Medicine website: ncbi.nlm.nih.gov/gorf/wblast2.cgi. Standard default parameter settings for UW-BLAST 2.0 are described at the following Internet site: blast.wustl.edu/blast/README.html#References. In addition, the BLAST algorithm uses the BLOSUM62 amino acid scoring matix, and optional parameters that can

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be used are as follows: (A) inclusion of a filter to mask segments of the query sequence that have low compositional complexity (as determined by the SEG program of Wootton and Federhen (Computers and Chemistry, 1993); also see Wootton and Federhen, 1996, Analysis of compositionally biased regions in sequence databases, *Methods Enzymol.* **266**: 554-71) or segments consisting of short-periodicity internal repeats (as determined by the XNU program of Claverie and States (Computers and Chemistry, 1993)), and (B) a statistical significance threshold for reporting matches against database sequences, or E-score (the expected probability of matches being found merely by chance, according to the stochastic model of Karlin and Altschul (1990); if the statistical significance ascribed to a match is greater than this E-score threshold, the match will not be reported.); preferred E-score threshold values are 0.5, or in order of increasing preference, 0.25, 0.1, 0.01, 0.001, 0.0001, 1e-10, 1e-20, 1e-30, 1e-40, 1e-50, 1e-75, or 1e-100.

"An isolated polypeptide consisting essentially of an amino acid sequence" means that the polypeptide may have, in addition to said amino acid sequence, additional material covalently linked to either or both ends of the polypeptide, said additional material preferably between 1 and 10,000 additional amino acids covalently linked to either end, each end, or both ends of polypeptide, and more preferably between 1 and 1,000 additional amino acids covalently linked to either end, each end, or both ends of the polypeptide, and most preferably between 1 and 100 additional amino acids covalently linked to either end, each end, or both ends of the polypeptide. In preferred embodiments, covalent linkage of additional amino acids to either end, each end, or both ends of the polypeptide results in a novel combined amino acid sequence that is neither naturally occurring nor disclosed in the art.

The present invention also provides for soluble forms of HPR1 and HPR2 polypeptides comprising or consisting essentially of certain fragments or domains of these polypeptides, and particularly those comprising the extracellular domain or one or more fragments of the extracellular domain. Soluble polypeptides are polypeptides that are capable of being secreted from the cells in which they are expressed. In such forms part or all of the intracellular and transmembrane domains of the polypeptide are deleted such that the polypeptide is fully secreted from the cell in which it is expressed. The intracellular and transmembrane domains of polypeptides of the invention can be identified in accordance with known techniques for determination of such domains from sequence information. Soluble HPR1 and HPR2 polypeptides also include those polypeptides which include part of the transmembrane region, provided that the soluble HPR1 or HPR2 polypeptide is capable of being secreted from a cell, and preferably retains HPR1 and/or HPR2 polypeptide activity. Soluble HPR1 and HPR2 polypeptides further include oligomers or fusion polypeptides comprising the extracellular portion of at least one HPR1 or HPR2 polypeptide, and fragments of any of these polypeptides that have HPR1 and/or HPR2 polypeptide activity. A secreted soluble polypeptide can be identified (and distinguished from its non-soluble membrane-bound counterparts) by separating intact cells which express the desired polypeptide from the culture medium, e.g., by centrifugation, and assaying the medium (supernatant) for the presence of the desired polypeptide. The presence of the desired polypeptide in the medium indicates that the polypeptide was secreted from the cells and thus is a

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soluble form of the polypeptide. The use of soluble forms of HPR1 or HPR2 polypeptides is advantageous for many applications. Purification of the polypeptides from recombinant host cells is facilitated, since the soluble polypeptides are secreted from the cells. Moreover, soluble polypeptides are generally more suitable than membrane-bound forms for parenteral administration and for many enzymatic procedures.

In another aspect of the invention, preferred polypeptides comprise various combinations of HPR1 and/or HPR2 polypeptide domains, such as the cytokine receptor domain and the intracellular signaling domain. Accordingly, polypeptides of the present invention and nucleic acids encoding them include those comprising or encoding two or more copies of a domain such as the cytokine receptor domain, two or more copies of a domain such as the intracellular signaling domain, or at least one copy of each domain, and these domains can be presented in any order within such polypeptides.

Further modifications in the peptide or DNA sequences can be made by those skilled in the art using known techniques. Modifications of interest in the polypeptide sequences can include the alteration, substitution, replacement, insertion or deletion of a selected amino acid. For example, one or more of the cysteine residues can be deleted or replaced with another amino acid to alter the conformation of the molecule, an alteration which may involve preventing formation of incorrect intramolecular disulfide bridges upon folding or renaturation. Techniques for such alteration, substitution, replacement, insertion or deletion are well known to those skilled in the art (see, e.g., U.S. Pat. No. 4,518,584). As another example, N-glycosylation sites in the polypeptide extracellular domain can be modified to preclude glycosylation, allowing expression of a reduced carbohydrate analog in mammalian and yeast expression systems. N-glycosylation sites in eukaryotic polypeptides are characterized by an amino acid triplet Asn-X-Y, wherein X is any amino acid except Pro and Y is Ser or Thr. Appropriate substitutions, additions, or deletions to the nucleotide sequence encoding these triplets will result in prevention of attachment of carbohydrate residues at the Asn side chain. Alteration of a single nucleotide, chosen so that Asn is replaced by a different amino acid, for example, is sufficient to inactivate an N-glycosylation site. Alternatively, the Ser or Thr can by replaced with another amino acid, such as Ala. Known procedures for inactivating N-glycosylation sites in polypeptides include those described in U.S. Patent 5,071,972 and EP 276,846. Additional variants within the scope of the invention include polypeptides that can be modified to create derivatives thereof by forming covalent or aggregative conjugates with other chemical moieties, such as glycosyl groups, lipids, phosphate, acetyl groups and the like. Covalent derivatives can be prepared by linking the chemical moieties to functional groups on amino acid side chains or at the N-terminus or C-terminus of a polypeptide. Conjugates comprising diagnostic (detectable) or therapeutic agents attached thereto are contemplated herein. Preferably, such alteration, substitution, replacement, insertion or deletion retains the desired activity of the polypeptide or a substantial equivalent thereof. One example is a variant that binds with essentially the same binding affinity as does the native form. Binding affinity can be measured by conventional procedures, e.g., as described in U.S. Patent No. 5,512,457 and as set forth herein.

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Other derivatives include covalent or aggregative conjugates of the polypeptides with other polypeptides or polypeptides, such as by synthesis in recombinant culture as N-terminal or C-terminal fusions. Examples of fusion polypeptides are discussed below in connection with oligomers. Further, fusion polypeptides can comprise peptides added to facilitate purification and identification. Such peptides include, for example, poly-His or the antigenic identification peptides described in U.S. Patent No. 5,011,912 and in Hopp et al., *Bio/Technology* 6:1204, 1988. One such peptide is the FLAG® peptide, which is highly antigenic and provides an epitope reversibly bound by a specific monoclonal antibody, enabling rapid assay and facile purification of expressed recombinant polypeptide. A murine hybridoma designated 4E11 produces a monoclonal antibody that binds the FLAG® peptide in the presence of certain divalent metal cations, as described in U.S. Patent 5,011,912. The 4E11 hybridoma cell line has been deposited with the American Type Culture Collection under accession no. HB 9259. Monoclonal antibodies that bind the FLAG® peptide are available from Eastman Kodak Co., Scientific Imaging Systems Division, New Haven, Connecticut.

Encompassed by the invention are oligomers or fusion polypeptides that contain an HPR1 polypeptide and/or an HPR2 polypeptide, one or more fragments of HPR1 and/or HPR2 polypeptides, or any of the derivative or variant forms of HPR1 and HPR2 polypeptides as disclosed herein. In particular embodiments, the oligomers comprise soluble HPR1 and/or HPR2 polypeptides. Oligomers can be in the form of covalently linked or non-covalently-linked multimers, including dimers, trimers, or higher oligomers. In one aspect of the invention, the oligomers maintain the binding ability of the polypeptide components and provide therefor, bivalent, trivalent, etc., binding sites. In an alternative embodiment the invention is directed to oligomers comprising multiple HPR1 and/or HPR2 polypeptides joined via covalent or non-covalent interactions between peptide moieties fused to the polypeptides, such peptides having the property of promoting oligomerization. Leucine zippers and certain polypeptides derived from antibodies are among the peptides that can promote oligomerization of the polypeptides attached thereto, as described in more detail below.

In embodiments where variants of the HPR1 and/or HPR2 polypeptides are constructed to include a membrane-spanning domain, they will form a Type I membrane polypeptide. Membrane-spanning HPR1 and/or HPR2 polypeptides can be fused with extracellular domains of receptor polypeptides for which the ligand is known. Such fusion polypeptides can then be manipulated to control the intracellular signaling pathways triggered by the membrane-spanning HPR1 or HPR2 polypeptide. HPR1 and HPR2 polypeptides that span the cell membrane can also be fused with agonists or antagonists of cell-surface receptors, or cellular adhesion molecules to further modulate HPR1 and/or HPR2 intracellular effects. In another aspect of the present invention, interleukins can be situated between the preferred HPR1 or HPR2 polypeptide fragment and other fusion polypeptide domains.

<u>Immunoglobulin-based Oligomers</u>. The polypeptides of the invention or fragments thereof can be fused to molecules such as immunoglobulins for many purposes, including increasing the valency of polypeptide binding sites. For example, fragments of an HPR1 polypeptide and/or of an

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HPR2 polypeptide can be fused directly or through linker sequences to the Fc portion of an immunoglobulin. For a bivalent form of the polypeptide, such a fusion could be to the Fc portion of an IgG molecule. Other immunoglobulin isotypes can also be used to generate such fusions. For example, a polypeptide-IgM fusion would generate a decavalent form of the polypeptide of the invention. The term "Fc polypeptide" as used herein includes native and mutein forms of polypeptides made up of the Fc region of an antibody comprising any or all of the CH domains of the Fc region. Truncated forms of such polypeptides containing the hinge region that promotes dimerization are also included. Preferred Fc polypeptides comprise an Fc polypeptide derived from a human IgG1 antibody. As one alternative, an oligomer is prepared using polypeptides derived from immunoglobulins. Preparation of fusion polypeptides comprising certain heterologous polypeptides fused to various portions of antibody-derived polypeptides (including the Fc domain) has been described, e.g., by Ashkenazi et al. (PNAS USA 88:10535, 1991); Byrn et al. (Nature 344:677, 1990); and Hollenbaugh and Aruffo ("Construction of Immunoglobulin Fusion Polypeptides", in Current Protocols in Immunology, Suppl. 4, pages 10.19.1 - 10.19.11, 1992). Methods for preparation and use of immunoglobulin-based oligomers are well known in the art. One embodiment of the present invention is directed to a dimer comprising two fusion polypeptides created by fusing a polypeptide of the invention to an Fc polypeptide derived from an antibody. A gene fusion encoding the polypeptide/Fc fusion polypeptide is inserted into an appropriate expression vector. Polypeptide/Fc fusion polypeptides are expressed in host cells transformed with the recombinant expression vector, and allowed to assemble much like antibody molecules, whereupon interchain disulfide bonds form between the Fc moieties to yield divalent molecules. One suitable Fc polypeptide, described in PCT application WO 93/10151, is a single chain polypeptide extending from the N-terminal hinge region to the native C-terminus of the Fc region of a human IgG1 antibody. Another useful Fc polypeptide is the Fc mutein described in U.S. Patent 5,457,035 and in Baum et al., (EMBO J. 13:3992-4001, 1994). The amino acid sequence of this mutein is identical to that of the native Fc sequence presented in WO 93/10151, except that amino acid 19 has been changed from Leu to Ala, amino acid 20 has been changed from Leu to Glu, and amino acid 22 has been changed from Gly to Ala. The mutein exhibits reduced affinity for Fc receptors. The above-described fusion polypeptides comprising Fc moieties (and oligomers formed therefrom) offer the advantage of facile purification by affinity chromatography over Polypeptide A or Polypeptide G columns. In other embodiments, the polypeptides of the invention can be substituted for the variable portion of an antibody heavy or light chain. If fusion polypeptides are made with both heavy and light chains of an antibody, it is possible to form an oligomer with as many as four HPR1 and/or HPR2 extracellular regions.

<u>Peptide-linker Based Oligomers.</u> Alternatively, the oligomer is a fusion polypeptide comprising multiple HPR1 and/or HPR2 polypeptides, with or without peptide linkers (spacer peptides). Among the suitable peptide linkers are those described in U.S. Patents 4,751,180 and 4,935,233. A DNA sequence encoding a desired peptide linker can be inserted between, and in the same reading frame as, the DNA sequences of the invention, using any suitable conventional technique.

For example, a chemically synthesized oligonucleotide encoding the linker can be ligated between the sequences. In particular embodiments, a fusion polypeptide comprises from two to four soluble HPR1 and/or HPR2 polypeptides, separated by peptide linkers. Suitable peptide linkers, their combination with other polypeptides, and their use are well known by those skilled in the art.

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Leucine-Zippers. Another method for preparing the oligomers of the invention involves use of a leucine zipper. Leucine zipper domains are peptides that promote oligomerization of the polypeptides in which they are found. Leucine zippers were originally identified in several DNA-binding polypeptides (Landschulz et al., *Science* 240:1759, 1988), and have since been found in a variety of different polypeptides. Among the known leucine zippers are naturally occurring peptides and derivatives thereof that dimerize or trimerize. The zipper domain (also referred to herein as an oligomerizing, or oligomer-forming, domain) comprises a repetitive heptad repeat, often with four or five leucine residues interspersed with other amino acids. Use of leucine zippers and preparation of oligomers using leucine zippers are well known in the art.

Other fragments and derivatives of the sequences of polypeptides which would be expected to retain polypeptide activity in whole or in part and may thus be useful for screening or other immunological methodologies can also be made by those skilled in the art given the disclosures herein. Such modifications are believed to be encompassed by the present invention.

### Nucleic Acids Encoding HPR1 Polypeptides and Nucleic Acids Encoding HPR2 Polypeptides

Encompassed within the invention are nucleic acids encoding HPR1 polypeptides and nucleic acids encoding HPR2 polypeptides. These nucleic acids can be identified in several ways, including isolation of genomic or cDNA molecules from a suitable source. Nucleotide sequences corresponding to the amino acid sequences described herein, to be used as probes or primers for the isolation of nucleic acids or as query sequences for database searches, can be obtained by "back-translation" from the amino acid sequences, or by identification of regions of amino acid identity with polypeptides for which the coding DNA sequence has been identified. The well-known polymerase chain reaction (PCR) procedure can be employed to isolate and amplify a DNA sequence encoding an HPR1 or HPR2 polypeptide or a desired combination of HPR1 and/or HPR2 polypeptide fragments. Oligonucleotides that define the desired termini of the combination of DNA fragments are employed as 5' and 3' primers. The oligonucleotides can additionally contain recognition sites for restriction endonucleases, to facilitate insertion of the amplified combination of DNA fragments into an expression vector. PCR techniques are described in Saiki et al., Science 239:487 (1988); Recombinant DNA Methodology, Wu et al., eds., Academic Press, Inc., San Diego (1989), pp. 189-196; and PCR Protocols: A Guide to Methods and Applications, Innis et. al., eds., Academic Press, Inc. (1990).

Nucleic acid molecules of the invention include DNA and RNA in both single-stranded and double-stranded form, as well as the corresponding complementary sequences. DNA includes, for example, cDNA, genomic DNA, chemically synthesized DNA, DNA amplified by PCR, and combinations thereof. The nucleic acid molecules of the invention include full-length genes or cDNA

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molecules as well as a combination of fragments thereof. The nucleic acids of the invention are preferentially derived from human sources, but the invention includes those derived from non-human species, as well.

An "isolated nucleic acid" is a nucleic acid that has been separated from adjacent genetic sequences present in the genome of the organism from which the nucleic acid was isolated, in the case of nucleic acids isolated from naturally-occurring sources. In the case of nucleic acids synthesized enzymatically from a template or chemically, such as PCR products, cDNA molecules, or oligonucleotides for example, it is understood that the nucleic acids resulting from such processes are isolated nucleic acids. An isolated nucleic acid molecule refers to a nucleic acid molecule in the form of a separate fragment or as a component of a larger nucleic acid construct. In one preferred embodiment, the invention relates to certain isolated nucleic acids that are substantially free from contaminating endogenous material. The nucleic acid molecule has preferably been derived from DNA or RNA isolated at least once in substantially pure form and in a quantity or concentration enabling identification, manipulation, and recovery of its component nucleotide sequences by standard biochemical methods (such as those outlined in Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, NY (1989)). Such sequences are preferably provided and/or constructed in the form of an open reading frame uninterrupted by internal non-translated sequences, or introns, that are typically present in eukaryotic genes. Sequences of non-translated DNA can be present 5' or 3' from an open reading frame, where the same do not interfere with manipulation or expression of the coding region.

"An isolated nucleic acid consisting essentially of a nucleotide sequence" means that the nucleic acid may have, in addition to said nucleotide sequence, additional material covalently linked to either or both ends of the nucleic acid molecule, said additional material preferably between 1 and 100,000 additional nucleotides covalently linked to either end, each end, or both ends of the nucleic acid molecule, and more preferably between 1 and 1,000 additional nucleotides covalently linked to either end, each end, or both ends of the nucleic acid molecule, and most preferably between 10 and 100 additional nucleotides covalently linked to either end, each end, or both ends of the nucleic acid molecule. In preferred embodiments, covalent linkage of additional nucleotides to either end, each end, or both ends of the nucleic acid molecule results in a novel combined nucleotide sequence that is neither naturally occurring nor disclosed in the art. An isolated nucleic acid consisting essentially of a nucleotide sequence may be an expression vector or other construct comprising said nucleotide sequence.

The present invention also includes nucleic acids that hybridize under moderately stringent conditions, and more preferably highly stringent conditions, to nucleic acids encoding HPR1 polypeptides and/or nucleic acids encoding HPR2 polypeptides described herein. The basic parameters affecting the choice of hybridization conditions and guidance for devising suitable conditions are set forth by Sambrook,, Fritsch, and Maniatis (1989, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., chapters 9 and 11; and Current Protocols

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in Molecular Biology, 1995, Ausubel et al., eds., John Wiley & Sons, Inc., sections 2.10 and 6.3-6.4), and can be readily determined by those having ordinary skill in the art based on, for example, the length and/or base composition of the DNA. One way of achieving moderately stringent conditions involves the use of a prewashing solution containing 5 x SSC, 0.5% SDS, 1.0 mM EDTA (pH 8.0), hybridization buffer of about 50% formamide, 6 x SSC, and a hybridization temperature of about 55 degrees C (or other similar hybridization solutions, such as one containing about 50% formamide, with a hybridization temperature of about 42 degrees C), and washing conditions of about 60 degrees C, in 0.5 x SSC, 0.1% SDS. Generally, highly stringent conditions are defined as hybridization conditions as above, but with washing at approximately 68 degrees C, 0.2 x SSC, 0.1% SDS. SSPE (1xSSPE is 0.15M NaCl, 10 mM NaH.sub.2 PO.sub.4, and 1.25 mM EDTA, pH 7.4) can be substituted for SSC (1xSSC is 0.15M NaCl and 15 mM sodium citrate) in the hybridization and wash buffers; washes are performed for 15 minutes after hybridization is complete. It should be understood that the wash temperature and wash salt concentration can be adjusted as necessary to achieve a desired degree of stringency by applying the basic principles that govern hybridization reactions and duplex stability, as known to those skilled in the art and described further below (see, e.g., Sambrook et al., 1989). When hybridizing a nucleic acid to a target nucleic acid of unknown sequence, the hybrid length is assumed to be that of the hybridizing nucleic acid. When nucleic acids of known sequence are hybridized, the hybrid length can be determined by aligning the sequences of the nucleic acids and identifying the region or regions of optimal sequence complementarity. The hybridization temperature for hybrids anticipated to be less than 50 base pairs in length should be 5 to 10.degrees C less than the melting temperature (Tm) of the hybrid, where Tm is determined according to the following equations. For hybrids less than 18 base pairs in length, Tm (degrees C) = 2(# of A + T bases) + 4(# of #G + C bases). For hybrids above 18 base pairs in length, Tm (degrees C) =  $81.5 + 16.6(\log_{10} [\text{Na}^+]) + 0.41(\% \text{ G} + \text{C})$ - (600/N), where N is the number of bases in the hybrid, and [Na<sup>+</sup>] is the concentration of sodium ions in the hybridization buffer ([Na<sup>+</sup>] for 1xSSC = 0.165M). Preferably, each such hybridizing nucleic acid has a length that is at least 15 nucleotides (or more preferably at least 18 nucleotides, or at least 20 nucleotides, or at least 25 nucleotides, or at least 30 nucleotides, or at least 40 nucleotides, or most preferably at least 50 nucleotides), or at least 25% (more preferably at least 50%, or at least 60%, or at least 70%, and most preferably at least 80%) of the length of the nucleic acid of the present invention to which it hybridizes, and has at least 60% sequence identity (more preferably at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, or at least 99%, and most preferably at least 99.5%) with the nucleic acid of the present invention to which it hybridizes, where sequence identity is determined by comparing the sequences of the hybridizing nucleic acids when aligned so as to maximize overlap and identity while minimizing sequence gaps as described in more detail above.

The present invention also provides genes corresponding to the nucleic acid sequences disclosed herein. "Corresponding genes" or "corresponding genomic nucleic acids" are the regions of the genome that are transcribed to produce the mRNAs from which cDNA nucleic acid sequences are

derived and can include contiguous regions of the genome necessary for the regulated expression of such genes. Corresponding genes can therefore include but are not limited to coding sequences, 5' and 3' untranslated regions, alternatively spliced exons, introns, promoters, enhancers, and silencer or suppressor elements. Corresponding genomic nucleic acids can include 10000 basepairs (more preferably, 5000 basepairs, still more preferably, 2500 basepairs, and most preferably, 1000 basepairs) of genomic nucleic acid sequence upstream of the first nucleotide of the genomic sequence corresponding to the initiation codon of the HPR1 coding sequence or of the HPR2 coding sequence, and 10000 basepairs (more preferably, 5000 basepairs, still more preferably, 2500 basepairs, and most preferably, 1000 basepairs) of genomic nucleic acid sequence downstream of the last nucleotide of the genomic sequence corresponding to the termination codon of the HPR1 coding sequence or of the HPR2 coding sequence. The corresponding genes or genomic nucleic acids can be isolated in accordance with known methods using the sequence information disclosed herein. Such methods include the preparation of probes or primers from the disclosed sequence information for identification and/or amplification of genes in appropriate genomic libraries or other sources of genomic materials. An "isolated gene" or an "isolated genomic nucleic acid" is a genomic nucleic acid that has been separated from the adjacent genomic sequences present in the genome of the organism from which the genomic nucleic acid was isolated.

### Methods for Making and Purifying HPR1 and HPR2 Polypeptides

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Methods for making HPR1 and HPR2 polypeptides are described below. Expression, isolation, and purification of the polypeptides and fragments of the invention can be accomplished by any suitable technique, including but not limited to the following methods. The isolated nucleic acid of the invention can be operably linked to an expression control sequence such as the pDC409 vector (Giri et al., 1990, EMBO J., 13: 2821) or the derivative pDC412 vector (Wiley et al., 1995, Immunity 3: 673). The pDC400 series vectors are useful for transient mammalian expression systems, such as CV-1 or 293 cells. Alternatively, the isolated nucleic acid of the invention can be linked to expression vectors such as pDC312, pDC316, or pDC317 vectors, which are useful for stable mammalian expression systems, such as CHO cells or their derivatives. Other expression control sequences and cloning technologies can also be used to produce the polypeptide recombinantly, such as the pMT2 or pED expression vectors (Kaufman et al., 1991, Nucleic Acids Res. 19: 4485-4490; and Pouwels et al., 1985, Cloning Vectors: A Laboratory Manual, Elsevier, New York) and the GATEWAY Vectors (lifetech.com/Content/Tech-Online/molecular\_biology/manuals\_pps/11797016.pdf; Life Technologies; Rockville, MD). Many suitable expression control sequences are known in the art. General methods of expressing recombinant polypeptides are also known and are exemplified in R. Kaufman, Methods in Enzymology 185, 537-566 (1990). As used herein "operably linked" means that the nucleic acid of the invention and an expression control sequence are situated within a construct, vector, or cell in such a way that the polypeptide encoded by the nucleic acid is expressed when appropriate molecules (such as polymerases) are present. As one embodiment of the invention, at least one expression control

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sequence is operably linked to the nucleic acid of the invention in a recombinant host cell or progeny thereof, the nucleic acid and/or expression control sequence having been introduced into the host cell by transformation or transfection, for example, or by any other suitable method. As another embodiment of the invention, at least one expression control sequence is integrated into the genome of a recombinant host cell such that it is operably linked to a nucleic acid sequence encoding a polypeptide of the invention. In a further embodiment of the invention, at least one expression control sequence is operably linked to a nucleic acid of the invention through the action of a trans-acting factor such as a transcription factor, either *in vitro* or in a recombinant host cell.

In addition, a sequence encoding an appropriate signal peptide (native or heterologous) can be incorporated into expression vectors. The choice of signal peptide or leader can depend on factors such as the type of host cells in which the recombinant polypeptide is to be produced. To illustrate, examples of heterologous signal peptides that are functional in mammalian host cells include the signal sequence for interleukin-7 (IL-7) described in United States Patent 4,965,195; the signal sequence for interleukin-2 receptor described in Cosman et al., Nature 312:768 (1984); the interleukin-4 receptor signal peptide described in EP 367,566; the type I interleukin-1 receptor signal peptide described in U.S. Patent 4,968,607; and the type II interleukin-1 receptor signal peptide described in EP 460,846. A DNA sequence for a signal peptide (secretory leader) can be fused in frame to the nucleic acid sequence of the invention so that the DNA is initially transcribed, and the mRNA translated, into a fusion polypeptide comprising the signal peptide. A signal peptide that is functional in the intended host cells promotes extracellular secretion of the polypeptide. The signal peptide is cleaved from the polypeptide upon secretion of polypeptide from the cell. The skilled artisan will also recognize that the position(s) at which the signal peptide is cleaved can differ from that predicted by computer program, and can vary according to such factors as the type of host cells employed in expressing a recombinant polypeptide. A polypeptide preparation can include a mixture of polypeptide molecules having different N-terminal amino acids, resulting from cleavage of the signal peptide at more than one site.

Established methods for introducing DNA into mammalian cells have been described (Kaufman, R.J., Large Scale Mammalian Cell Culture, 1990, pp. 15-69). Additional protocols using commercially available reagents, such as Lipofectamine lipid reagent (Gibco/BRL) or Lipofectamine-Plus lipid reagent, can be used to transfect cells (Felgner et al., Proc. Natl. Acad. Sci. USA 84:7413-7417, 1987). In addition, electroporation can be used to transfect mammalian cells using conventional procedures, such as those in Sambrook et al. (Molecular Cloning: A Laboratory Manual, 2 ed. Vol. 1-3, Cold Spring Harbor Laboratory Press, 1989). Selection of stable transformants can be performed using methods known in the art, such as, for example, resistance to cytotoxic drugs. Kaufman et al., Meth. in Enzymology 185:487-511, 1990, describes several selection schemes, such as dihydrofolate reductase (DHFR) resistance. A suitable strain for DHFR selection can be CHO strain DX-B11, which is deficient in DHFR (Urlaub and Chasin, Proc. Natl. Acad. Sci. USA 77:4216-4220, 1980). Other examples of selectable markers that can be incorporated into an expression vector include cDNAs

conferring resistance to antibiotics, such as G418 and hygromycin B. Cells harboring the vector can be selected on the basis of resistance to these compounds.

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Alternatively, gene products can be obtained via homologous recombination, or "gene targeting," techniques. Such techniques employ the introduction of exogenous transcription control elements (such as the CMV promoter or the like) in a particular predetermined site on the genome, to induce expression of the endogenous nucleic acid sequence of interest (see, for example, U.S. Patent No. 5,272,071). The location of integration into a host chromosome or genome can be easily determined by one of skill in the art, given the known location and sequence of the gene. In a preferred embodiment, the present invention also contemplates the introduction of exogenous transcriptional control elements in conjunction with an amplifiable gene, to produce increased amounts of the gene product, again, without the need for isolation of the gene sequence itself from the host cell.

A number of types of cells can act as suitable host cells for expression of the polypeptide. Mammalian host cells include, for example, the COS-7 line of monkey kidney cells (ATCC CRL 1651) (Gluzman et al., Cell 23:175, 1981), L cells, C127 cells, 3T3 cells (ATCC CCL 163), Chinese hamster ovary (CHO) cells, HeLa cells, BHK (ATCC CRL 10) cell lines, the CV1/EBNA cell line derived from the African green monkey kidney cell line CV1 (ATCC CCL 70) as described by McMahan et al. (EMBO J. 10: 2821, 1991), human kidney 293 cells, human epidermal A431 cells, human Colo205 cells, other transformed primate cell lines, normal diploid cells, cell strains derived from in vitro culture of primary tissue, primary explants, HL-60, U937, HaK or Jurkat cells. Alternatively, it is possible to produce the polypeptide in lower eukaryotes such as yeast or in prokaryotes such as bacteria. Potentially suitable yeasts include Saccharomyces cerevisiae, Schizosaccharomyces pombe, Kluyveromyces strains, Candida, or any yeast strain capable of expressing heterologous polypeptides. Potentially suitable bacterial strains include Escherichia coli, Bacillus subtilis, Salmonella typhimurium, or any bacterial strain capable of expressing heterologous polypeptides. If the polypeptide is made in yeast or bacteria, it may be necessary to modify the polypeptide produced therein, for example by phosphorylation or glycosylation of the appropriate sites, in order to obtain the functional polypeptide. Such covalent attachments can be accomplished using known chemical or enzymatic methods. The polypeptide can also be produced by operably linking the isolated nucleic acid of the invention to suitable control sequences in one or more insect expression vectors, and employing an insect expression system. Materials and methods for baculovirus/insect cell expression systems are commercially available in kit form from, e.g., Invitrogen, San Diego, Calif., U.S.A. (the MaxBac® kit), and such methods are well known in the art, as described in Summers and Smith, Texas Agricultural Experiment Station Bulletin No. 1555 (1987), and Luckow and Summers, Bio/Technology 6:47 (1988). Cell-free translation systems could also be employed to produce polypeptides using RNAs derived from nucleic acid constructs disclosed herein. A host cell that comprises an isolated nucleic acid of the invention, preferably operably linked to at least one expression control sequence, is a "recombinant host cell".

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The polypeptide of the invention can be prepared by culturing transformed host cells under The resulting expressed culture conditions suitable to express the recombinant polypeptide. polypeptide can then be purified from such culture (i.e., from culture medium or cell extracts) using known purification processes, such as gel filtration and ion exchange chromatography. The purification of the polypeptide can also include an affinity column containing agents which will bind to the polypeptide; one or more column steps over such affinity resins as concanavalin A-agarose, heparintoyopearl® or Cibacrom blue 3GA Sepharose®; one or more steps involving hydrophobic interaction chromatography using such resins as phenyl ether, butyl ether, or propyl ether; or immunoaffinity chromatography. Alternatively, the polypeptide of the invention can also be expressed in a form which will facilitate purification. For example, it can be expressed as a fusion polypeptide, such as those of maltose binding polypeptide (MBP), glutathione-S-transferase (GST) or thioredoxin (TRX). Kits for expression and purification of such fusion polypeptides are commercially available from New England BioLab (Beverly, Mass.), Pharmacia (Piscataway, N.J.) and InVitrogen, respectively. The polypeptide can also be tagged with an epitope and subsequently purified by using a specific antibody directed to such epitope. One such epitope (FLAG®) is commercially available from Kodak (New Haven, Conn.). Finally, one or more reverse-phase high performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media, e.g., silica gel having pendant methyl or other aliphatic groups, can be employed to further purify the polypeptide. Some or all of the foregoing purification steps, in various combinations, can also be employed to provide a substantially homogeneous isolated recombinant polypeptide. The polypeptide thus purified is substantially free of other mammalian polypeptides and is defined in accordance with the present invention as an "isolated polypeptide"; such isolated polypeptides of the invention include isolated antibodies that bind to HPR1 and/or HPR2 polypeptides, fragments, variants, binding partners etc. The polypeptide of the invention can also be expressed as a product of transgenic animals, e.g., as a component of the milk of transgenic cows, goats, pigs, or sheep which are characterized by somatic or germ cells containing a nucleotide sequence encoding the polypeptide.

It is also possible to utilize an affinity column comprising a polypeptide-binding polypeptide of the invention, such as a monoclonal antibody generated against polypeptides of the invention, to affinity-purify expressed polypeptides. These polypeptides can be removed from an affinity column using conventional techniques, e.g., in a high salt elution buffer and then dialyzed into a lower salt buffer for use or by changing pH or other components depending on the affinity matrix utilized, or be competitively removed using the naturally occurring substrate of the affinity moiety, such as a polypeptide derived from the invention. In this aspect of the invention, polypeptide-binding polypeptides, such as the anti-polypeptide antibodies of the invention or other polypeptides that can interact with the polypeptide of the invention, can be bound to a solid phase support such as a column chromatography matrix or a similar substrate suitable for identifying, separating, or purifying cells that express polypeptides of the invention on their surface. Adherence of polypeptide-binding polypeptides of the invention to a solid phase contacting surface can be accomplished by any means, for example,

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magnetic microspheres can be coated with these polypeptide-binding polypeptides and held in the incubation vessel through a magnetic field. Suspensions of cell mixtures are contacted with the solid phase that has such polypeptide-binding polypeptides thereon. Cells having polypeptides of the invention on their surface bind to the fixed polypeptide-binding polypeptide and unbound cells then are washed away. This affinity-binding method is useful for purifying, screening, or separating such polypeptide-expressing cells from solution. Methods of releasing positively selected cells from the solid phase are known in the art and encompass, for example, the use of enzymes. Such enzymes are preferably non-toxic and non-injurious to the cells and are preferably directed to cleaving the cell-surface binding partner. Alternatively, mixtures of cells suspected of containing polypeptide-expressing cells of the invention first can be incubated with a biotinylated polypeptide-binding polypeptide of the invention. The resulting mixture then is passed through a column packed with avidin-coated beads, whereby the high affinity of biotin for avidin provides the binding of the polypeptide-binding cells to the beads. Use of avidin-coated beads is known in the art. See Berenson, et al. *J. Cell. Biochem.*, 10D:239 (1986). Wash of unbound material and the release of the bound cells is performed using conventional methods.

The polypeptide can also be produced by known conventional chemical synthesis. The synthetically-constructed polypeptide sequences, by virtue of sharing primary, secondary or tertiary structural and/or conformational characteristics with HPR1 and/or HPR2 polypeptides can possess biological properties in common therewith, including HPR1 and/or HPR2 polypeptide activity. Thus, they can be employed as biologically active or immunological substitutes for natural, purified polypeptides in screening of therapeutic compounds and in immunological processes for the development of antibodies.

The desired degree of purity depends on the intended use of the polypeptide. A relatively high degree of purity is desired when the polypeptide is to be administered *in vivo*, for example. In such a case, the polypeptides are purified such that no polypeptide bands corresponding to other polypeptides are detectable upon analysis by SDS-polyacrylamide gel electrophoresis (SDS-PAGE). It will be recognized by one skilled in the pertinent field that multiple bands corresponding to the polypeptide can be visualized by SDS-PAGE, due to differential glycosylation, differential post-translational processing, and the like. Most preferably, the polypeptide of the invention is purified to substantial homogeneity, as indicated by a single polypeptide band upon analysis by SDS-PAGE. The polypeptide band can be visualized by silver staining, Coomassie blue staining, or (if the polypeptide is radiolabeled) by autoradiography.

# Antagonists and Agonists of HPR1 and/or HPR2 Polypeptides

Any method which neutralizes HPR1 and/or HPR2 polypeptides or inhibits expression of the HPR1 and/or HPR2 genes (either transcription or translation) can be used to reduce the biological activities of HPR1 and/or HPR2 polypeptides. In particular embodiments, antagonists inhibit the binding of at least one HPR1 polypeptide and/or at least one HPR2 polypeptide to cells, thereby inhibiting biological activities induced by the binding of those HPR1 or HPR2 polypeptides to the

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cells. In certain other embodiments of the invention, antagonists can be designed to reduce the level of endogenous HPR1 and/or HPR2 gene expression, e.g., using well-known antisense or ribozyme approaches to inhibit or prevent translation of HPR1 and/or HPR2 mRNA transcripts; triple helix approaches to inhibit transcription of HPR1 and/or HPR2 genes; or targeted homologous recombination to inactivate or "knock out" the HPR1 gene(s), the HPR2 gene(s), or their endogenous promoters or enhancer elements. Such antisense, ribozyme, and triple helix antagonists can be designed to reduce or inhibit either unimpaired, or if appropriate, mutant HPR1 and/or HPR2 gene activity. Techniques for the production and use of such molecules are well known to those of skill in the art.

Antisense RNA and DNA molecules act to directly block the translation of mRNA by hybridizing to targeted mRNA and preventing polypeptide translation. Antisense approaches involve the design of oligonucleotides (either DNA or RNA) that are complementary to an HPR1 and/or to an HPR2 mRNA. The antisense oligonucleotides will bind to the complementary target gene mRNA transcripts and prevent translation. Absolute complementarity, although preferred, is not required. A sequence "complementary" to a portion of a nucleic acid, as referred to herein, means a sequence having sufficient complementarity to be able to hybridize with the nucleic acid, forming a stable duplex (or triplex, as appropriate). In the case of double-stranded antisense nucleic acids, a single strand of the duplex DNA can thus be tested, or triplex formation can be assayed. The ability to hybridize will depend on both the degree of complementarity and the length of the antisense nucleic acid. Preferred oligonucleotides are complementary to the 5' end of the message, e.g., the 5' untranslated sequence up to and including the AUG initiation codon. However, oligonucleotides complementary to the 5'- or 3'non- translated, non-coding regions of the HPR1 or HPR2 gene transcript(s) could be used in an antisense approach to inhibit translation of endogenous HPR1 and/or HPR2 mRNA. Antisense nucleic acids should be at least six nucleotides in length, and are preferably oligonucleotides ranging from 6 to about 50 nucleotides in length. In specific aspects the oligonucleotide is at least 10 nucleotides, at least 17 nucleotides, at least 25 nucleotides or at least 50 nucleotides. The oligonucleotides can be DNA or RNA or chimeric mixtures or derivatives or modified versions thereof, single-stranded or doublestranded. Chimeric oligonucleotides, oligonucleosides, or mixed oligonucleotides/oligonucleosides of the invention can be of several different types. These include a first type wherein the "gap" segment of nucleotides is positioned between 5' and 3' "wing" segments of linked nucleosides and a second "open end" type wherein the "gap" segment is located at either the 3' or the 5' terminus of the oligomeric compound (see, e.g., U.S. Pat. No. 5,985,664). Oligonucleotides of the first type are also known in the art as "gapmers" or gapped oligonucleotides. Oligonucleotides of the second type are also known in the art as "hemimers" or "wingmers". The oligonucleotide can be modified at the base moiety, sugar moiety, or phosphate backbone, for example, to improve stability of the molecule, hybridization, etc. The oligonucleotide can include other appended groups such as peptides (e.g., for targeting host cell receptors in vivo), or agents facilitating transport across the cell membrane (see, e.g., Letsinger et al., 1989, Proc Natl Acad Sci USA 86:6553-6556; Lemaitre et al., 1987, Proc Natl Acad Sci 84:648-652;

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PCT Publication No. WO88/09810), or hybridization-triggered cleavage agents or intercalating agents. (See, e.g., Zon, 1988, Pharm. Res. 5:539-549). The antisense molecules should be delivered to cells which express the HPR1 and/or HPR2 transcript in vivo. A number of methods have been developed for delivering antisense DNA or RNA to cells; e.g., antisense molecules can be injected directly into the tissue or cell derivation site, or modified antisense molecules, designed to target the desired cells (e.g., antisense linked to peptides or antibodies that specifically bind receptors or antigens expressed on the target cell surface) can be administered systemically. However, it is often difficult to achieve intracellular concentrations of the antisense sufficient to suppress translation of endogenous mRNAs. Therefore a preferred approach utilizes a recombinant DNA construct in which the antisense oligonucleotide is placed under the control of a strong pol III or pol II promoter. The use of such a construct to transfect target cells in the patient will result in the transcription of sufficient amounts of single stranded RNAs that will form complementary base pairs with the endogenous HPR1 and/or HPR2 gene transcripts and thereby prevent translation of the HPR1 and/or HPR2 mRNA. For example, a vector can be introduced in vivo such that it is taken up by a cell and directs the transcription of an antisense RNA. Such a vector can remain episomal or become chromosomally integrated, as long as it can be transcribed to produce the desired antisense RNA. Such vectors can be constructed by recombinant DNA technology methods standard in the art. Vectors can be plasmid, viral, or others known in the art, used for replication and expression in mammalian cells.

Ribozyme molecules designed to catalytically cleave HPR1 and/or HPR2 mRNA transcripts can also be used to prevent translation of HPR1 and/or HPR2 mRNA and expression of HPR1 and/or HPR2 polypeptides. (See, e.g., PCT International Publication WO90/11364 and US Patent No. 5,824,519). The ribozymes that can be used in the present invention include hammerhead ribozymes (Haseloff and Gerlach, 1988, Nature, 334:585-591), RNA endoribonucleases (hereinafter "Cech-type ribozymes") such as the one which occurs naturally in Tetrahymena Thermophila (known as the IVS, or L-19 IVS RNA) and which has been extensively described by Thomas Cech and collaborators (International Patent Application No. WO 88/04300; Been and Cech, 1986, Cell, 47:207-216). As in the antisense approach, the ribozymes can be composed of modified oligonucleotides (e.g. for improved stability, targeting, etc.) and should be delivered to cells which express HPR1 and/or HPR2 polypeptides in vivo. A preferred method of delivery involves using a DNA construct "encoding" the ribozyme under the control of a strong constitutive pol III or pol II promoter, so that transfected cells will produce sufficient quantities of the ribozyme to destroy endogenous HPR1 and/or HPR2 messages and inhibit translation. Because ribozymes, unlike antisense molecules, are catalytic, a lower intracellular concentration is required for efficiency.

Alternatively, endogenous HPR1 and/or HPR2 gene expression can be reduced by targeting deoxyribonucleotide sequences complementary to the regulatory region of the target gene (i.e., the target gene promoter and/or enhancers) to form triple helical structures that prevent transcription of the target HPR1 and/or HPR2 gene. (See generally, Helene, 1991, Anticancer Drug Des., 6(6), 569-584; Helene, et al., 1992, Ann. N.Y. Acad. Sci., 660, 27-36; and Maher, 1992, Bioassays 14(12), 807-815).

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Anti-sense RNA and DNA, ribozyme, and triple helix molecules of the invention can be prepared by any method known in the art for the synthesis of DNA and RNA molecules. These include techniques for chemically synthesizing oligodeoxyribonucleotides and oligoribonucleotides well known in the art such as for example solid phase phosphoramidite chemical synthesis. Oligonucleotides can be synthesized by standard methods known in the art, e.g. by use of an automated DNA synthesizer (such as are commercially available from Biosearch, Applied Biosystems, etc.). As examples, phosphorothioate oligonucleotides can be synthesized by the method of Stein et al., 1988, Nucl. Acids Res. 16:3209. Methylphosphonate oligonucleotides can be prepared by use of controlled pore glass polymer supports (Sarin et al., 1988, Proc. Natl. Acad. Sci. U.S.A. 85:7448-7451). Alternatively, RNA molecules can be generated by in vitro and in vivo transcription of DNA sequences encoding the antisense RNA molecule. Such DNA sequences can be incorporated into a wide variety of vectors that incorporate suitable RNA polymerase promoters such as the T7 or SP6 polymerase promoters. Alternatively, antisense cDNA constructs that synthesize antisense RNA constitutively or inducibly, depending on the promoter used, can be introduced stably into cell lines.

Endogenous target gene expression can also be reduced by inactivating or "knocking out" the target gene or its promoter using targeted homologous recombination (e.g., see Smithies, et al., 1985, Nature 317, 230-234; Thomas and Capecchi, 1987, Cell 51, 503-512; Thompson, et al., 1989, Cell 5, 313-321). For example, a mutant, non-functional target gene (or a completely unrelated DNA sequence) flanked by DNA homologous to the endogenous target gene (either the coding regions or regulatory regions of the target gene) can be used, with or without a selectable marker and/or a negative selectable marker, to transfect cells that express the target gene in vivo. Insertion of the DNA construct, via targeted homologous recombination, results in inactivation of the target gene. Such approaches are particularly suited in the agricultural field where modifications to ES (embryonic stem) cells can be used to generate animal offspring with an inactive target gene (e.g., see Thomas and Capecchi, 1987 and Thompson, 1989, supra), or in model organisms such as Caenorhabditis elegans where the "RNA interference" ("RNAi") technique (Grishok, Tabara, and Mello, 2000, Genetic requirements for inheritance of RNAi in C. elegans, Science 287 (5462): 2494-2497), or the introduction of transgenes (Dernburg et al., 2000, Transgene-mediated cosuppression in the C. elegans germ line, Genes Dev. 14 (13): 1578-1583) are used to inhibit the expression of specific target genes. However this approach can be adapted for use in humans provided the recombinant DNA constructs are directly administered or targeted to the required site in vivo using appropriate vectors such as viral vectors.

Organisms that have enhanced, reduced, or modified expression of the gene(s) corresponding to the nucleic acid sequences disclosed herein are provided. The desired change in gene expression can be achieved through the use of antisense nucleic acids or ribozymes that bind and/or cleave the mRNA transcribed from the gene (Albert and Morris, 1994, Trends Pharmacol. Sci. 15(7): 250-254; Lavarosky et al., 1997, Biochem. Mol. Med. 62(1): 11-22; and Hampel, 1998, Prog. Nucleic Acid Res. Mol. Biol. 58: 1-39). Transgenic animals that have multiple copies of the gene(s) corresponding to the nucleic

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acid sequences disclosed herein, preferably produced by transformation of cells with genetic constructs that are stably maintained within the transformed cells and their progeny, are provided. Transgenic animals that have modified genetic control regions that increase or reduce gene expression levels, or that change temporal or spatial patterns of gene expression, are also provided (see European Patent No. 0 649 464 B1). In addition, organisms are provided in which the gene(s) corresponding to the nucleic acid sequences disclosed herein have been partially or completely inactivated, through insertion of extraneous sequences into the corresponding gene(s) or through deletion of all or part of the corresponding gene(s). Partial or complete gene inactivation can be accomplished through insertion, preferably followed by imprecise excision, of transposable elements (Plasterk, 1992, Bioessays 14(9): 629-633; Zwaal et al., 1993, Proc Natl Acad Sci USA 90(16): 7431-7435; Clark et al., 1994, Proc Natl Acad Sci USA 91(2): 719-722), or through homologous recombination, preferably detected by positive/negative genetic selection strategies (Mansour et al., 1988, Nature 336: 348-352; U.S. Pat. Nos. 5,464,764; 5,487,992; 5,627,059; 5,631,153; 5,614,396; 5,616,491; and 5,679,523). These organisms with altered gene expression are preferably eukaryotes and more preferably are mammals. Such organisms are useful for the development of non-human models for the study of disorders involving the corresponding gene(s), and for the development of assay systems for the identification of molecules that interact with the polypeptide product(s) of the corresponding gene(s).

Also encompassed within the invention are HPR1 and HPR2 polypeptide variants with partner binding sites that have been altered in conformation so that (1) the HPR1 or HPR2 variant will still bind to its partner(s), but a specified small molecule will fit into the altered binding site and block that interaction, or (2) the HPR1 or HPR2 variant will no longer bind to its partner(s) unless a specified small molecule is present (see for example Bishop *et al.*, 2000, *Nature* 407: 395-401). Nucleic acids encoding such altered HPR1 or HPR2 polypeptides can be introduced into organisms according to methods described herein, and can replace the endogenous nucleic acid sequences encoding the corresponding HPR1 or HPR2 polypeptide. Such methods allow for the interaction of a particular HPR1 or HPR2 polypeptide with its binding partners to be regulated by administration of a small molecule compound to an organism, either systemically or in a localized manner.

The HPR1 and HPR2 polypeptides themselves can also be employed in inhibiting a biological activity of HPR1 and /or of HPR2 in *in vitro* or *in vivo* procedures. Encompassed within the invention are cytokine receptor domains of HPR1 and HPR2 polypeptides that act as "dominant negative" inhibitors of native HPR1 and/or HPR2 polypeptide function when expressed as fragments or as components of fusion polypeptides. For example, a purified polypeptide domain of the present invention can be used to inhibit binding of HPR1 or HPR2 polypeptides to endogenous binding partners. Such use effectively would block HPR1 and/or HPR2 polypeptide interactions and inhibit HPR1 and/or HPR2 polypeptide activities. In still another aspect of the invention, a soluble form of an HPR1 and/or HPR2 binding partner is used to bind to an endogenous HPR1 and/or HPR2 polypeptide, and competitively inhibit activation of that endogenous HPR1 and/or HPR2 polypeptide. Furthermore, antibodies which bind to HPR1 and/or HPR2 polypeptides often inhibit HPR1 and/or HPR2

polypeptide activity and act as antagonists. For example, antibodies that specifically recognize one or more epitopes of HPR1 and/or HPR2 polypeptides, or epitopes of conserved variants of HPR1 and/or HPR2 polypeptides, or peptide fragments of an HPR1 and/or HPR2 polypeptide can be used in the invention to inhibit HPR1 and/or HPR2 polypeptide activity. Such antibodies include but are not limited to polyclonal antibodies, monoclonal antibodies (mAbs), humanized or chimeric antibodies, single chain antibodies, Fab fragments, F(ab')2 fragments, fragments produced by a Fab expression library, anti-idiotypic (anti-Id) antibodies, and epitope-binding fragments of any of the above. Alternatively, purified and modified HPR1 and/or HPR2 polypeptides of the present invention can be administered to modulate interactions between HPR1 and/or HPR2 polypeptides and HPR1 and/or HPR2 binding partners that are not membrane-bound. Such an approach will allow an alternative method for the modification of HPR1- and/or HPR2-influenced bioactivity.

In an alternative aspect, the invention further encompasses the use of agonists of HPR1 and/or HPR2 polypeptide activity to treat or ameliorate the symptoms of a disease for which increased HPR1 and/or HPR2 polypeptide activity is beneficial. Such diseases include but are not limited to pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, osteoporosis resulting from a lack of bone-forming cells, obesity, deficient mammary development, and infertility. In a preferred aspect, the invention entails administering compositions comprising an HPR1 or HPR2 nucleic acid or an HPR1 or HPR2 polypeptide to cells in vitro, to cells ex vivo, to cells in vivo, and/or to a multicellular organism such as a vertebrate or mammal. Preferred therapeutic forms of HPR1 and HPR2 are soluble forms, as described above. In still another aspect of the invention, the compositions comprise administering an HPR1-encoding nucleic acid or an HPR2-encoding nucleic acid for expression of an HPR1 or HPR2 polypeptide in a host organism for treatment of disease. Particularly preferred in this regard is expression in a human patient for treatment of a dysfunction associated with aberrant (e.g., decreased) endogenous activity of an HPR1 or HPR2 polypeptide. Furthermore, the invention encompasses the administration to cells and/or organisms of compounds found to increase the endogenous activity of HPR1 and/or HPR2 polypeptides. One example of compounds that increase HPR1 and/or HPR2 polypeptide activity are agonistic antibodies, preferably monoclonal antibodies, that bind to HPR1 and/or HPR2 polypeptides or binding partners, which may increase HPR1 and/or HPR2 polypeptide activity by causing constitutive intracellular signaling (or "ligand mimicking"), or by preventing the binding of a native inhibitor of HPR1 and/or HPR2 polypeptide activity.

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#### Antibodies to HPR1 and/or HPR2 Polypeptides

Antibodies that are immunoreactive with the polypeptides of the invention are provided herein. Such antibodies specifically bind to the polypeptides via the antigen-binding sites of the antibody (as opposed to non-specific binding). In the present invention, specifically binding antibodies are those that will specifically recognize and bind with HPR1 and/or HPR2 polypeptides, homologues, and variants, but not with other molecules. In one preferred embodiment, the antibodies are specific for

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the polypeptides of the present invention and do not cross-react with other polypeptides. In this manner, the HPR1 and HPR2 polypeptides, fragments, variants, fusion polypeptides, etc., as set forth above can be employed as "immunogens" in producing antibodies immunoreactive therewith.

More specifically, the polypeptides, fragment, variants, fusion polypeptides, etc. contain antigenic determinants or epitopes that elicit the formation of antibodies. These antigenic determinants or epitopes can be either linear or conformational (discontinuous). Linear epitopes are composed of a single section of amino acids of the polypeptide, while conformational or discontinuous epitopes are composed of amino acids sections from different regions of the polypeptide chain that are brought into close proximity upon polypeptide folding (Janeway and Travers, Immuno Biology 3:9 (Garland Publishing Inc., 2nd ed. 1996)). Because folded polypeptides have complex surfaces, the number of epitopes available is quite numerous; however, due to the conformation of the polypeptide and steric hinderances, the number of antibodies that actually bind to the epitopes is less than the number of available epitopes (Janeway and Travers, Immuno Biology 2:14 (Garland Publishing Inc., 2nd ed. 1996)). Epitopes can be identified by any of the methods known in the art. Thus, one aspect of the present invention relates to the antigenic epitopes of the polypeptides of the invention. Such epitopes are useful for raising antibodies, in particular monoclonal antibodies, as described in more detail below. Additionally, epitopes from the polypeptides of the invention can be used as research reagents, in assays, and to purify specific binding antibodies from substances such as polyclonal sera or supernatants from cultured hybridomas. Such epitopes or variants thereof can be produced using techniques well known in the art such as solid-phase synthesis, chemical or enzymatic cleavage of a polypeptide, or using recombinant DNA technology.

As to the antibodies that can be elicited by the epitopes of the polypeptides of the invention, whether the epitopes have been isolated or remain part of the polypeptides, both polyclonal and monoclonal antibodies can be prepared by conventional techniques. See, for example, Monoclonal Antibodies, Hybridomas: A New Dimension in Biological Analyses, Kennet et al. (eds.), Plenum Press, New York (1980); and Antibodies: A Laboratory Manual, Harlow and Land (eds.), Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, (1988); Kohler and Milstein, (U.S. Pat. No. 4,376,110); the human B-cell hybridoma technique (Kosbor et al., 1984, J Immunol 133: 3001-3005; Cole et al., 1983, Proc Natl Acad Sci USA 80:2026-2030); and the EBV-hybridoma technique (Cole et al., 1985, Monoclonal Antibodies And Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). Hybridoma cell lines that produce monoclonal antibodies specific for the polypeptides of the invention are also contemplated herein. Such hybridomas can be produced and identified by conventional techniques. The hybridoma producing the mAb of this invention can be cultivated in vitro or in vivo. Production of high titers of mAbs in vivo makes this the presently preferred method of production. One method for producing such a hybridoma cell line comprises immunizing an animal with a polypeptide; harvesting spleen cells from the immunized animal; fusing said spleen cells to a myeloma cell line, thereby generating hybridoma cells; and identifying a hybridoma cell line that produces a monoclonal antibody that binds the polypeptide. For the production of antibodies, various host animals can be immunized by

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injection with one or more of the following: an HPR1 or HPR2 polypeptide, a fragment of an HPR1 or HPR2 polypeptide, a functional equivalent of an HPR1 or HPR2 polypeptide, or a mutant form of an HPR1 or HPR2 polypeptide. Such host animals can include but are not limited to rabbits, mice, and rats. Various adjuvants can be used to increase the immunologic response, depending on the host species, including but not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanin, dinitrophenol, and potentially useful human adjutants such as BCG (bacille Calmette-Guerin) and Corynebacterium parvum. The monoclonal antibodies can be recovered by conventional techniques. Such monoclonal antibodies can be of any immunoglobulin class including IgG, IgM, IgE, IgA, IgD and any subclass thereof.

In addition, techniques developed for the production of "chimeric antibodies" (Takeda et al., 1985, Nature, 314: 452-454; Morrison et al., 1984, Proc Natl Acad Sci USA 81: 6851-6855; Boulianne et al., 1984, Nature 312: 643-646; Neuberger et al., 1985, Nature 314: 268-270) by splicing the genes from a mouse antibody molecule of appropriate antigen specificity together with genes from a human antibody molecule of appropriate biological activity can be used. A chimeric antibody is a molecule in which different portions are derived from different animal species, such as those having a variable region derived from a porcine mAb and a human immunoglobulin constant region. The monoclonal antibodies of the present invention also include humanized versions of murine monoclonal antibodies. Such humanized antibodies can be prepared by known techniques and offer the advantage of reduced immunogenicity when the antibodies are administered to humans. In one embodiment, a humanized monoclonal antibody comprises the variable region of a murine antibody (or just the antigen binding site thereof) and a constant region derived from a human antibody. Alternatively, a humanized antibody fragment can comprise the antigen binding site of a murine monoclonal antibody and a variable region fragment (lacking the antigen-binding site) derived from a human antibody. Procedures for the production of chimeric and further engineered monoclonal antibodies include those described in Riechmann et al. (Nature 332:323, 1988), Liu et al. (PNAS 84:3439, 1987), Larrick et al. (Bio/Technology 7:934, 1989), and Winter and Harris (TIPS 14:139, Can, 1993). Useful techniques for humanizing antibodies are also discussed in U.S. Patent 6,054,297. Procedures to generate antibodies transgenically can be found in GB 2,272,440, US Patent Nos. 5,569,825 and 5,545,806, and related patents. Preferably, for use in humans, the antibodies are human or humanized; techniques for creating such human or humanized antibodies are also well known and are commercially available from, for example, Medarex Inc. (Princeton, NJ) and Abgenix Inc. (Fremont, CA). In another preferred embodiment, fully human antibodies for use in humans are produced by screening a phage display library of human antibody variable domains (Vaughan et al., 1998, Nat Biotechnol. 16(6): 535-539; and U.S. Patent No. 5,969,108).

Antigen-binding antibody fragments which recognize specific epitopes can be generated by known techniques. For example, such fragments include but are not limited to: the F(ab')2 fragments which can be produced by pepsin digestion of the antibody molecule and the Fab fragments which can

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be generated by reducing the disulfide bridges of the (ab')2 fragments. Alternatively, Fab expression libraries can be constructed (Huse *et al.*, 1989, Science, 246:1275-1281) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity. Techniques described for the production of single chain antibodies (U.S. Pat. No. 4,946,778; Bird, 1988, Science 242:423-426; Huston *et al.*, 1988, Proc. Natl. Acad. Sci. USA 85:5879-5883; and Ward *et al.*, 1989, Nature 334:544-546) can also be adapted to produce single chain antibodies against HPR1 and/or HPR2 gene products. Single chain antibodies are formed by linking the heavy and light chain fragments of the Fv region via an amino acid bridge, resulting in a single chain polypeptide. Such single chain antibodies can also be useful intracellularly (i.e., as 'intrabodies), for example as described by Marasco *et al.* (*J. Immunol. Methods* 231:223-238, 1999) for genetic therapy in HIV infection. In addition, antibodies to the HPR1 and/or HPR2 polypeptide can, in turn, be utilized to generate anti-idiotype antibodies that "mimic" the HPR1 and/or HPR2 polypeptide and that may bind to the binding partner(s) of HPR1 and/or HPR2 polypeptides, using techniques well known to those skilled in the art. (See, *e.g.*, Greenspan & Bona, 1993, FASEB J 7(5):437-444; and Nissinoff, 1991, J. Immunol. 147(8):2429-2438).

Antibodies that are immunoreactive with the polypeptides of the invention include bispecific antibodies (i.e., antibodies that are immunoreactive with the polypeptides of the invention via a first antigen binding domain, and also immunoreactive with a different polypeptide via a second antigen binding domain). A variety of bispecific antibodies have been prepared, and found useful both in vitro and in vivo (see, for example, U.S. Patent 5,807,706; and Cao and Suresh, 1998, Bioconjugate Chem 9: 635-644). Numerous methods of preparing bispecific antibodies are known in the art, including the use of hybrid-hybridomas such as quadromas, which are formed by fusing two differed hybridomas, and triomas, which are formed by fusing a hybridoma with a lymphocyte (Milstein and Cuello, 1983, Nature 305: 537-540; U.S. Patent 4,474,893; and U.S. Patent 6,106,833). U.S. Patent 6,060,285 discloses a process for the production of bispecific antibodies in which at least the genes for the light chain and the variable portion of the heavy chain of an antibody having a first specificity are transfected into a hybridoma cell secreting an antibody having a second specificity. Chemical coupling of antibody fragments has also been used to prepare antigen-binding molecules having specificity for two different antigens (Brennan et al., 1985, Science 229: 81-83; Glennie et al., J. Immunol., 1987, 139:2367-2375; and U.S. Patent 6,010,902). Bispecific antibodies can also be produced via recombinant means, for example, by using, the leucine zipper moieties from the Fos and Jun proteins (which preferentially form heterodimers) as described by Kostelny et al. (J. Immnol. 148:1547-4553; 1992). U.S. Patent 5,582,996 discloses the use of complementary interactive domains (such as leucine zipper moieties or other lock and key interactive domain structures) to facilitate heterodimer formation in the production of bispecific antibodies. Tetravalent, bispecific molecules can be prepared by fusion of DNA encoding the heavy chain of an F(ab')2 fragment of an antibody with either DNA encoding the heavy chain of a second F(ab')2 molecule (in which the CH1 domain is replaced by a CH3 domain), or with DNA encoding a single chain FV fragment of an antibody, as described in U.S. Patent 5,959,083. Expression of the resultant fusion genes in mammalian cells, together with the genes for the

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corresponding light chains, yields tetravalent bispecific molecules having specificity for selected antigens. Bispecific antibodies can also be produced as described in U.S. Patent 5,807,706. Generally, the method involves introducing a protuberance (constructed by replacing small amino acid side chains with larger side chains) at the interface of a first polypeptide and a corresponding cavity (prepared by replacing large amino acid side chains with smaller ones) in the interface of a second polypeptide. Moreover, single-chain variable fragments (sFvs) have been prepared by covalently joining two variable domains; the resulting antibody fragments can form dimers or trimers, depending on the length of a flexible linker between the two variable domains (Kortt *et al.*, 1997, *Protein Engineering* 10:423-433).

Screening procedures by which such antibodies can be identified are well known, and can involve immunoaffinity chromatography, for example. Antibodies can be screened for agonistic (i.e., ligand-mimicking) properties. Such antibodies, upon binding to cell surface HPR1 and/or HPR2, induce biological effects (e.g., transduction of biological signals) similar to the biological effects induced when the HPR1 and/or HPR2 binding partner binds to cell surface HPR1 and/or HPR2. Agonistic antibodies can be used to induce HPR1- and/or HPR2-mediated intracellular signaling or cell proliferation. Bispecific antibodies can be identified by screening with two separate assays, or with an assay wherein the bispecific antibody serves as a bridge between the first antigen and the second antigen (the latter is coupled to a detectable moiety). Bispecific antibodies that bind HPR1 and/or HPR2 polypeptides of the invention via a first antigen binding domain will be useful in diagnostic applications and in treating cell proliferation, differentiation, or activation diseases or conditions. Examples of polypeptides (or other antigens) that the inventive bispecific antibodies bind via a second antigen binding domain include: four alpha helix bundle cytokines such as IL-6, OSM, LIF, CNTF, CLC, IL-12p35, and IL-23p19; soluble hematopoietin receptors such as EBI-3, soluble IL-6R alpha, cytokine-like factor-1 (CLF), IL-12p40, or a soluble form of HPR1 and/or HPR2; and soluble hematopoietin receptors such as EBI-3 etc. in conjunction with a four alpha helix bundle cytokine.

Those antibodies that can block binding of the HPR1 and/or HPR2 polypeptides of the invention to binding partners for HPR1 and/or HPR2 can be used to inhibit HPR1- and/or HPR2-mediated intracellular signaling or cell proliferation that results from such binding. Such blocking antibodies can be identified using any suitable assay procedure, such as by testing antibodies for the ability to inhibit binding of HPR1 and/or HPR2 to certain cells expressing an HPR1 and/or HPR2 binding partner. Alternatively, blocking antibodies can be identified in assays for the ability to inhibit a biological effect that results from binding of soluble HPR1 and/or HPR2 to target cells. Antibodies can be assayed for the ability to inhibit HPR1 and/or HPR2 binding partner-mediated cell stimulatory pathways, for example. Such an antibody can be employed in an *in vitro* procedure, or administered *in vivo* to inhibit a biological activity mediated by the entity that generated the antibody. Disorders caused or exacerbated (directly or indirectly) by the interaction of HPR1 and/or HPR2 with cell surface binding partner receptor thus can be treated. A therapeutic method involves *in vivo* administration of a blocking antibody to a mammal in an amount effective in inhibiting HPR1 and/or HPR2 binding

partner-mediated biological activity. Monoclonal antibodies are generally preferred for use in such therapeutic methods. In one embodiment, an antigen-binding antibody fragment is employed. Compositions comprising an antibody that is directed against HPR1 and/or HPR2, and a physiologically acceptable diluent, excipient, or carrier, are provided herein. Suitable components of such compositions are as described below for compositions containing HPR1 and/or HPR2 polypeptides.

Also provided herein are conjugates comprising a detectable (e.g., diagnostic) or therapeutic agent, attached to the antibody. Examples of such agents are presented above. The conjugates find use in *in vitro* or *in vivo* procedures. The antibodies of the invention can also be used in assays to detect the presence of the polypeptides or fragments of the invention, either *in vitro* or *in vivo*. The antibodies also can be employed in purifying polypeptides or fragments of the invention by immunoaffinity chromatography.

### Rational Design of Compounds that Interact with HPR1 and/or HPR2 Polypeptides

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The goal of rational drug design is to produce structural analogs of biologically active polypeptides of interest or of small molecules with which they interact, e.g., inhibitors, agonists, antagonists, etc. Any of these examples can be used to fashion drugs which are more active or stable forms of the polypeptide or which enhance or interfere with the function of a polypeptide in vivo (Hodgson J (1991) Biotechnology 9:19-21). In one approach, the three-dimensional structure of a polypeptide of interest, or of a polypeptide-inhibitor complex, is determined by x-ray crystallography, by nuclear magnetic resonance, or by computer homology modeling or, most typically, by a combination of these approaches. Both the shape and charges of the polypeptide must be ascertained to elucidate the structure and to determine active site(s) of the molecule. Less often, useful information regarding the structure of a polypeptide may be gained by modeling based on the structure of homologous polypeptides. In both cases, relevant structural information is used to design analogous HPR1- and/or HPR2-like molecules, to identify efficient inhibitors, or to identify small molecules that bind HPR1 and/or HPR2 polypeptides. Useful examples of rational drug design include molecules which have improved activity or stability as shown by Braxton S and Wells JA (1992 Biochemistry 31:7796-7801) or which act as inhibitors, agonists, or antagonists of native peptides as shown by Athauda SB et al (1993 J Biochem 113:742-746). The use of HPR1 and/or HPR2 polypeptide structural information in molecular modeling software systems to assist in inhibitor design and in studying inhibitor-HPR1 polypeptide and/or inhibitor-HPR2 polypeptide interaction is also encompassed by the invention. A particular method of the invention comprises analyzing the threedimensional structure of HPR1 and/or HPR2 polypeptides for likely binding sites of substrates, synthesizing a new molecule that incorporates a predictive reactive site, and assaying the new molecule as described further herein.

It is also possible to isolate a target-specific antibody, selected by functional assay, as described further herein, and then to solve its crystal structure. This approach, in principle, yields a pharmacore upon which subsequent drug design can be based. It is possible to bypass polypeptide

crystallography altogether by generating anti-idiotypic antibodies (anti-ids) to a functional, pharmacologically active antibody. As a mirror image of a mirror image, the binding site of the anti-ids would be expected to be an analog of the original receptor. The anti-id could then be used to identify and isolate peptides from banks of chemically or biologically produced peptides. The isolated peptides would then act as the pharmacore.

## 10 Assays of HPR1 and HPR2 Polypeptide Activities

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The purified HPR1 and HPR2 polypeptides of the invention (including polypeptides, polypeptides, fragments, variants, oligomers, and other forms) are useful in a variety of assays. For example, the HPR1 and HPR2 molecules of the present invention can be used to identify binding partners of HPR1 and/or HPR2 polypeptides, which can also be used to modulate intracellular signaling, cell proliferation, or immune cell activity. Alternatively, they can be used to identify non-binding-partner molecules or substances that modulate intracellular signaling, cell proliferation, or immune cell activity.

Assays to Identify Binding Partners. HPR1 and HPR2 polypeptides and fragments thereof can be used to identify binding partners. For example, they can be tested for the ability to bind a candidate binding partner in any suitable assay, such as a conventional binding assay. To illustrate, the HPR1 or HPR2 polypeptide can be labeled with a detectable reagent (e.g., a radionuclide, chromophore, enzyme that catalyzes a colorimetric or fluorometric reaction, and the like). The labeled polypeptide is contacted with cells expressing the candidate binding partner. The cells then are washed to remove unbound labeled polypeptide, and the presence of cell-bound label is determined by a suitable technique, chosen according to the nature of the label.

One example of a binding assay procedure is as follows. A recombinant expression vector containing the candidate binding partner cDNA is constructed. CV1-EBNA-1 cells in 10 cm2 dishes are transfected with this recombinant expression vector. CV-1/EBNA-1 cells (ATCC CRL 10478) constitutively express EBV nuclear antigen-1 driven from the CMV Immediate-early enhancer/promoter. CV1-EBNA-1 was derived from the African Green Monkey kidney cell line CV-1 (ATCC CCL 70), as described by McMahan et al., (EMBO J. 10:2821, 1991). The transfected cells are cultured for 24 hours, and the cells in each dish then are split into a 24-well plate. After culturing an additional 48 hours, the transfected cells (about 4 x 10<sup>4</sup> cells/well) are washed with BM-NFDM, which is binding medium (RPMI 1640 containing 25 mg/ml bovine serum albumin, 2 mg/ml sodium azide, 20 mM Hepes pH 7.2) to which 50 mg/ml nonfat dry milk has been added. The cells then are incubated for 1 hour at 37°C with various concentrations of, for example, a soluble polypeptide/Fc fusion polypeptide made as set forth above. Cells then are washed and incubated with a constant saturating concentration of a 125I-mouse anti-human IgG in binding medium, with gentle agitation for 1 hour at 37°C. After extensive washing, cells are released via trypsinization. The mouse anti-human IgG employed above is directed against the Fc region of human IgG and can be obtained from Jackson Immunoresearch Laboratories, Inc., West Grove, PA. The antibody is radioiodinated using the standard chloramine-T method. The antibody will bind to the Fc portion of any polypeptide/Fc

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polypeptide that has bound to the cells. In all assays, non-specific binding of <sup>125</sup>I-antibody is assayed in the absence of the Fc fusion polypeptide/Fc, as well as in the presence of the Fc fusion polypeptide and a 200-fold molar excess of unlabeled mouse anti-human IgG antibody. Cell-bound 125I-antibody is quantified on a Packard Autogamma counter. Affinity calculations (Scatchard, Ann. N.Y. Acad. Sci. 51:660, 1949) are generated on RS/1 (BBN Software, Boston, MA) run on a Microvax computer. Binding can also be detected using methods that are well suited for high-throughput screening procedures, such as scintillation proximity assays (Udenfriend et al., 1985, Proc Natl Acad Sci USA 82: 8672-8676), homogeneous time-resolved fluorescence methods (Park et al., 1999, Anal Biochem 269: 94-104), fluorescence resonance energy transfer (FRET) methods (Clegg RM, 1995, Curr Opin Biotechnol 6: 103-110), or methods that measure any changes in surface plasmon resonance when a bound polypeptide is exposed to a potential binding partner, using for example a biosensor such as that supplied by Biacore AB (Uppsala, Sweden). Compounds that can be assayed for binding to HPR1 and/or HPR2 polypeptides include but are not limited to small organic molecules, such as those that are commercially available - often as part of large combinatorial chemistry compound 'libraries' - from companies such as Sigma-Aldrich (St. Louis, MO), Arqule (Woburn, MA), Enzymed (Iowa City, IA), Maybridge Chemical Co.(Trevillett, Cornwall, UK), MDS Panlabs (Bothell, WA), Pharmacopeia (Princeton, NJ), and Trega (San Diego, CA). Preferred small organic molecules for screening using these assays are usually less than 10K molecular weight and can possess a number of physicochemical and pharmacological properties which enhance cell penetration, resist degradation, and/or prolong their physiological half-lives (Gibbs, J., 1994, Pharmaceutical Research in Molecular Oncology, Cell 79(2): 193-198). Compounds including natural products, inorganic chemicals, and biologically active materials such as proteins and toxins can also be assayed using these methods for the ability to bind to HPR1 and/or HPR2 polypeptides.

Yeast Two-Hybrid or "Interaction Trap" Assays. Because HPR1 and HPR2 polypeptides bind or potentially bind to another polypeptide (such as, for example, in a receptor-ligand interaction), the nucleic acid encoding the HPR1 or HPR2 polypeptide can also be used in interaction trap assays (such as, for example, that described in Gyuris et al., Cell 75:791-803 (1993)) to identify nucleic acids encoding the other polypeptide with which binding occurs, or to identify inhibitors of the binding interaction. Polypeptides involved in these binding interactions can also be used to screen for peptide or small molecule inhibitors or agonists of the binding interaction.

Competitive Binding Assays. Another type of suitable binding assay is a competitive binding assay. To illustrate, biological activity of a variant can be determined by assaying for the variant's ability to compete with the native polypeptide for binding to the candidate binding partner. Competitive binding assays can be performed by conventional methodology. Reagents that can be employed in competitive binding assays include radiolabeled HPR1 or HPR2 and intact cells expressing HPR1 and/or HPR2 (endogenous or recombinant) on the cell surface. For example, a radiolabeled soluble HPR1 or HPR2 fragment can be used to compete with a soluble HPR1 variant and/or a soluble HPR2 variant for binding to cell surface receptors. Instead of intact cells, one could

substitute a soluble binding partner/Fc fusion polypeptide bound to a solid phase through the interaction of Polypeptide A or Polypeptide G (on the solid phase) with the Fc moiety. Chromatography columns that contain Polypeptide A and Polypeptide G include those available from Pharmacia Biotech, Inc., Piscataway, NJ.

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Assays to Identify Modulators of Intracellular Signaling, Cell Proliferation, or Immune Cell Activity. The influence of HPR1 or HPR2 on intracellular signaling, cell proliferation, or immune cell activity can be manipulated to control these activities in target cells. For example, the disclosed HPR1 and HPR2 polypeptides, nucleic acids encoding the disclosed HPR1 and HPR2 polypeptides, or agonists or antagonists of such polypeptides can be administered to a cell or group of cells to induce, enhance, suppress, or arrest intracellular signaling or cell proliferation by the target cells. Identification of HPR1 and HPR2 polypeptides, agonists or antagonists that can be used in this manner can be carried out via a variety of assays known to those skilled in the art. Included in such assays are those that evaluate the ability of an HPR1 or HPR2 polypeptide to influence intracellular signaling, cell proliferation, or immune cell activity. Such an assay would involve, for example, the analysis of immune cell interaction in the presence of an HPR1 polypeptide and/or an HPR1 polypeptide. In such an assay, one would determine a rate of intracellular signaling or cell proliferation in the presence of the HPR1 and/or HPR2 polypeptide and then determine if such intracellular signaling or cell proliferation is altered in the presence of a candidate agonist or antagonist or another HPR1 or HPR2 polypeptide. Exemplary assays for this aspect of the invention include cytokine secretion assays, cell proliferation assays, and mixed lymphocyte reactions involving antigen presenting cells and T cells. These assays are well known to those skilled in the art.

In another aspect, the present invention provides a method of detecting the ability of a test compound to affect the intracellular signaling or cell proliferation activity of a cell. In this aspect, the method comprises: (1) contacting a first group of target cells with a test compound including an HPR1 polypeptide and/or an HPR2 polypeptide, or a fragment or fragments thereof, under conditions appropriate to the particular assay being used; (2) measuring the net rate of intracellular signaling or cell proliferation among the target cells; and (3) observing the net rate of intracellular signaling or cell proliferation among control cells contacting the HPR1 and./or HPR2 polypeptides or fragments thereof, in the absence of a test compound, under otherwise identical conditions as the first group of cells. In this embodiment, the net rate of intracellular signaling or cell proliferation in the control cells is compared to that of the cells treated with both a test compound and the HPR1 and/or HPR2 polypeptide(s). The comparison will provide a difference in the net rate of intracellular signaling or cell proliferation can be identified. The test compound can function as an effector by either activating or up-regulating, or by inhibiting or down-regulating, intracellular signaling or cell proliferation, and can be detected through this method.

<u>Cell Proliferation, Cell Death, Cell Differentiation, and Cell Adhesion Assays</u>. A polypeptide of the present invention may exhibit cytokine, cell proliferation (either inducing or inhibiting), or cell differentiation (either inducing or inhibiting) activity, or may induce production of other cytokines in

certain cell populations. Many polypeptide factors discovered to date have exhibited such activity in one or more factor-dependent cell proliferation assays, and hence the assays serve as a convenient confirmation of cell stimulatory activity. The activity of a polypeptide of the present invention is evidenced by any one of a number of routine factor-dependent cell proliferation assays for cell lines including, without limitation, 32D, DA2, DA1G, T10, B9, B9/11, BaF3, MC9/G, M+ (preB M+), 2E8, RB5, DA1, 123, T1165, HT2, CTLL2, TF-1, Mo7e and CMK. The activity of an HPR1 or HPR2 polypeptide of the invention may, among other means, be measured by the following methods:

Assays for T-cell or thymocyte proliferation include without limitation those described in: Current Protocols in Immunology, Coligan et al. eds, Greene Publishing Associates and Wiley-Interscience (pp. 3.1-3.19: In vitro assays for mouse lymphocyte function; Chapter 7: Immunologic studies in humans); Takai et al., J. Immunol. 137: 3494-3500, 1986; Bertagnolli et al., J. Immunol. 145: 1706-1712, 1990; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Bertagnolli, et al., J. Immunol. 149:3778-3783, 1992; Bowman et al., J. Immunol. 152: 1756-1761, 1994.

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Assays for cytokine production and/or proliferation of spleen cells, lymph node cells or thymocytes include, without limitation, those described in: Kruisbeek and Shevach, 1994, Polyclonal T cell stimulation, in *Current Protocols in Immunology*, Coligan *et al.* eds. Vol 1 pp. 3.12.1-3.12.14, John Wiley and Sons, Toronto; and Schreiber, 1994, Measurement of mouse and human interferon gamma in *Current Protocols in Immunology*, Coligan *et al.* eds. Vol 1 pp. 6.8.1-6.8.8, John Wiley and Sons, Toronto.

Assays for proliferation and differentiation of hematopoietic and lymphopoietic cells include, without limitation, those described in: Bottomly et al., 1991, Measurement of human and murine interleukin 2 and interleukin 4, in Current Protocols in Immunology, Coligan et al. eds. Vol 1 pp. 6.3.1-6.3.12, John Wiley and Sons, Toronto; deVries et al., J Exp Med 173: 1205-1211, 1991; Moreau et al., Nature 336:690-692, 1988; Greenberger et al., Proc Natl Acad Sci.USA 80: 2931-2938, 1983; Nordan, 1991, Measurement of mouse and human interleukin 6, in Current Protocols in Immunology Coligan et al. eds. Vol 1 pp. 6.6.1-6.6.5, John Wiley and Sons, Toronto; Smith et al., Proc Natl Acad Sci USA 83: 1857-1861, 1986; Bennett et al., 1991, Measurement of human interleukin 11, in Current Protocols in Immunology Coligan et al. eds. Vol 1 pp. 6.15.1 John Wiley and Sons, Toronto; Ciarletta et al., 1991, Measurement of mouse and human Interleukin 9, in Current Protocols in Immunology Coligan et al. eds. Vol 1 pp. 6.13.1, John Wiley and Sons, Toronto.

Assays for T-cell clone responses to antigens (which will identify, among others, polypeptides that affect APC-T cell interactions as well as direct T-cell effects by measuring proliferation and cytokine production) include, without limitation, those described in: *Current Protocols in Immunology*, Coligan *et al.* eds, Greene Publishing Associates and Wiley-Interscience (Chapter 3: *In vitro* assays for mouse lymphocyte function; Chapter 6: Cytokines and their cellular receptors; Chapter 7: Immunologic studies in humans); Weinberger et al., Proc Natl Acad Sci USA 77: 6091-6095, 1980; Weinberger et al., Eur. J. Immun. 11:405-411, 1981; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988

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Assays for thymocyte or splenocyte cytotoxicity include, without limitation, those described in: Current Protocols in Immunology, Coligan et al. eds, Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Herrmann et al., Proc. Natl. Acad. Sci. USA 78:2488-2492, 1981; Herrmann et al., J. Immunol. 128:1968-1974, 1982; Handa et al., J. Immunol. 135:1564-1572, 1985; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Herrmann et al., Proc. Natl. Acad. Sci. USA 78:2488-2492, 1981; Herrmann et al., J. Immunol. 128:1968-1974, 1982; Handa et al., J. Immunol. 135:1564-1572, 1985; Takai et al., J. Immunol. 137:3494-3500, 1986; Bowmanet al., J. Virology 61:1992-1998; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Brown et al., J. Immunol. 153:3079-3092, 1994.

Assays for T-cell-dependent immunoglobulin responses and isotype switching (which will identify, among others, polypeptides that modulate T-cell dependent antibody responses and that affect Th1/Th2 profiles) include, without limitation, those described in: Maliszewski, J Immunol 144: 3028-3033, 1990; and Mond and Brunswick, 1994, Assays for B cell function: *in vitro* antibody production, in *Current Protocols in Immunology* Coligan *et al.* eds. Vol 1 pp. 3.8.1-3.8.16, John Wiley and Sons, Toronto.

Mixed lymphocyte reaction (MLR) assays (which will identify, among others, polypeptides that generate predominantly Th1 and CTL responses) include, without limitation, those described in: Current Protocols in Immunology, Coligan et al. eds, Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., J. Immunol. 149:3778-3783, 1992.

Dendritic cell-dependent assays (which will identify, among others, polypeptides expressed by dendritic cells that activate naive T-cells) include, without limitation, those described in: Guery et al., J. Immunol 134:536-544, 1995; Inaba et al., J Exp Med 173:549-559, 1991; Macatonia et al., J Immunol 154:5071-5079, 1995; Porgador et al., J Exp Med 182:255-260, 1995; Nair et al., J Virology 67:4062-4069, 1993; Huang et al., Science 264:961-965, 1994; Macatonia et al., J Exp Med 169:1255-1264, 1989; Bhardwaj et al., J Clin Invest 94:797-807, 1994; and Inaba et al., J Exp Med 172:631-640,1990.

Assays for lymphocyte survival/apoptosis (which will identify, among others, polypeptides that prevent apoptosis after superantigen induction and polypeptides that regulate lymphocyte homeostasis) include, without limitation, those described in: Darzynkiewicz et al., Cytometry 13:795-808, 1992; Gorczyca et al., Leukemia 7:659-670, 1993; Gorczyca et al., Cancer Research 53:1945-1951, 1993; Itoh et al., Cell 66:233-243, 1991; Zacharchuk, J Immunol 145:4037-4045, 1990; Zamai et al., Cytometry 14:891-897, 1993; Gorczyca et al., International Journal of Oncology 1:639-648, 1992.

Assays for polypeptides that influence early steps of T-cell commitment and development include, without limitation, those described in: Antica et al., Blood 84:111-117, 1994; Fine et al., Cell Immunol 155:111-122, 1994; Galy et al., Blood 85:2770-2778, 1995; Toki et al., Proc Natl Acad Sci. USA 88:7548-7551, 1991

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Assays for embryonic stem cell differentiation (which will identify, among others, polypeptides that influence embryonic differentiation hematopoiesis) include, without limitation, those described in: Johansson et al. Cellular Biology 15:141-151, 1995; Keller et al., Molecular and Cellular Biology 13:473-486, 1993; McClanahan et al., Blood 81:2903-2915, 1993.

Assays for stem cell survival and differentiation (which will identify, among others, polypeptides that regulate lympho-hematopoiesis) include, without limitation, those described in: Methylcellulose colony forming assays, Freshney, 1994, In *Culture of Hematopoietic Cells*, Freshney et al. eds. pp. 265-268, Wiley-Liss, Inc., New York, NY; Hirayama et al., Proc. Natl. Acad. Sci. USA 89:5907-5911, 1992; Primitive hematopoietic colony forming cells with high proliferative potential, McNiece and Briddell, 1994, In *Culture of Hematopoietic Cells*, Freshney et al. eds. pp. 23-39, Wiley-Liss, Inc., New York, NY; Neben et al., Experimental Hematology 22:353-359, 1994; Ploemacher, 1994, Cobblestone area forming cell assay, In *Culture of Hematopoietic Cells*, Freshney et al. eds. pp. 1-21, Wiley-Liss, Inc., New York, NY; Spooncer et al., 1994, Long term bone marrow cultures in the presence of stromal cells, In *Culture of Hematopoietic Cells*, Freshney et al. eds. pp. 163-179, Wiley-Liss, Inc., New York, NY; Sutherland, 1994, Long term culture initiating cell assay, In *Culture of Hematopoietic Cells*, Freshney et al. eds. Vol pp. 139-162, Wiley-Liss, Inc., New York, NY.

Assay for hemostatic and thrombolytic activity include, without limitation, those described in: Linet et al., J. Clin. Pharmacol. 26:131-140, 1986; Burdick et al., Thrombosis Res. 45:413-419,1987; Humphrey et al., Fibrinolysis 5:71-79 (1991); Schaub, Prostaglandins 35:467-474, 1988.

Assays for receptor-ligand activity include without limitation those described in: *Current Protocols in Immunology* Coligan *et al.* eds, Greene Publishing Associates and Wiley-Interscience (Chapter 7.28, Measurement of cellular adhesion under static conditions 7.28.1-7.28.22), Takai et al., Proc. Natl. Acad. Sci. USA 84:6864-6868, 1987; Bierer et al., J. Exp. Med. 168:1145-1156, 1988; Rosenstein et al., J. Exp. Med. 169:149-160 1989; Stoltenborg et al., J. Immunol. Methods 175:59-68, 1994; Stitt et al., Cell 80:661-670, 1995.

#### 30 Diagnostic and Other Uses of HPR1 and HPR2 Polypeptides and Nucleic Acids

The nucleic acids encoding the HPR1 and HPR2 polypeptides provided by the present invention can be used for numerous diagnostic or other useful purposes. The nucleic acids of the invention can be used to express recombinant polypeptide for analysis, characterization or therapeutic use; as markers for tissues in which the corresponding polypeptide is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in disease states); as molecular weight markers on Southern gels; as chromosome markers or tags (when labeled) to identify chromosomes or to map related gene positions; to compare with endogenous DNA sequences in patients to identify potential genetic disorders; as probes to hybridize and thus discover novel, related DNA sequences; as a source of information to derive PCR primers for genetic fingerprinting; as a probe to "subtract-out" known sequences in the process of discovering other novel nucleic acids; for selecting and making oligomers for attachment to a "gene chip" or other support, including for examination of expression patterns; to raise anti-polypeptide antibodies using DNA immunization

techniques; as an antigen to raise anti-DNA antibodies or elicit another immune response, and. for gene therapy. Uses of HPR1 and HPR2 polypeptides and fragmented polypeptides include, but are not limited to, the following: purifying polypeptides and measuring the activity thereof; delivery agents; therapeutic and research reagents; molecular weight and isoelectric focusing markers; controls for peptide fragmentation; identification of unknown polypeptides; and preparation of antibodies. Any or all nucleic acids suitable for these uses are capable of being developed into reagent grade or kit format for commercialization as products. Methods for performing the uses listed above are well known to those skilled in the art. References disclosing such methods include without limitation "Molecular Cloning: A Laboratory Manual", 2d ed., Cold Spring Harbor Laboratory Press, Sambrook, J., E. F. Fritsch and T. Maniatis eds., 1989, and "Methods in Enzymology: Guide to Molecular Cloning Techniques", Academic Press, Berger, S. L. and A. R. Kimmel eds., 1987

Probes and Primers. Among the uses of the disclosed HPR1 and HPR2 nucleic acids, and combinations of fragments thereof, is the use of fragments as probes or primers. Such fragments generally comprise at least about 17 contiguous nucleotides of a DNA sequence. In other embodiments, a DNA fragment comprises at least 30, or at least 60, contiguous nucleotides of a DNA sequence. The basic parameters affecting the choice of hybridization conditions and guidance for devising suitable conditions are set forth by Sambrook et al., 1989 and are described in detail above. Using knowledge of the genetic code in combination with the amino acid sequences set forth above, sets of degenerate oligonucleotides can be prepared. Such oligonucleotides are useful as primers, e.g., in polymerase chain reactions (PCR), whereby DNA fragments are isolated and amplified. In certain embodiments, degenerate primers can be used as probes for non-human genetic libraries. Such libraries would include but are not limited to cDNA libraries, genomic libraries, and even electronic EST (express sequence tag) or DNA libraries. Homologous sequences identified by this method would then be used as probes to identify non-human HPR1 and HPR2 homologues.

Chromosome Mapping. The nucleic acids encoding HPR1 and HPR2 polypeptides, and the disclosed fragments and combinations of these nucleic acids, can be used by those skilled in the art using well-known techniques to identify the human chromosome to which these nucleic acids map. Useful techniques include, but are not limited to, using the sequence or portions, including oligonucleotides, as a probe in various well-known techniques such as radiation hybrid mapping (high resolution), in situ hybridization to chromosome spreads (moderate resolution), and Southern blot hybridization to hybrid cell lines containing individual human chromosomes (low resolution). Alternatively, the genomic sequences corresponding to nucleic acids encoding a cytokine polypeptide of the invention are mapped by comparison to sequences in public and proprietary databases, such as GenBank (ncbi.nlm.nih.gov/BLAST), Locuslink (ncbi.nlm.nih.gov:80/LocusLink/), Unigene (ncbi.nlm.nih.gov/cgi-bin/UniGene), AceView (ncbi.nlm.nih.gov/AceView), Gene Map Viewer (ncbi.nlm.nih.gov/genemap), Online Mendelian Inheritance in Man (OMIM) (ncbi.nlm.nih.gov/Omim), and proprietary databases such as the Celera Discovery System (celera.com). These computer analyses of available genomic sequence information can provide the identification of the specific chromosomal

location of human and/or murine genomic sequences corresponding to sequences encoding HPR1 or HPR2 polypeptides of the invention, and the unique genetic mapping relationships between HPR1 or HPR2 genomic sequences and the genetic map locations of known human genetic disorders

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<u>Diagnostics and Gene Therapy.</u> The nucleic acids encoding HPR1 and HPR2 polypeptides, and the disclosed fragments and combinations of these nucleic acids can be used by one skilled in the art using well-known techniques to analyze abnormalities associated with the genes corresponding to these polypeptides. This enables one to distinguish conditions in which this marker is rearranged or deleted. In addition, nucleic acids of the invention or a fragment thereof can be used as a positional marker to map other genes of unknown location. The DNA can be used in developing treatments for any disorder mediated (directly or indirectly) by defective, or insufficient amounts of, the genes corresponding to the nucleic acids of the invention. Disclosure herein of native nucleotide sequences permits the detection of defective genes, and the replacement thereof with normal genes. Defective genes can be detected in *in vitro* diagnostic assays, and by comparison of a native nucleotide sequence disclosed herein with that of a gene derived from a person suspected of harboring a defect in this gene.

Methods of Screening for Binding Partners. The HPR1 and HPR2 polypeptides of the invention each can be used as reagents in methods to screen for or identify binding partners. For example, the HPR1 and HPR2 polypeptides can be attached to a solid support material and may bind to their binding partners in a manner similar to affinity chromatography. In particular embodiments, a polypeptide is attached to a solid support by conventional procedures. As one example, chromatography columns containing functional groups that will react with functional groups on amino acid side chains of polypeptides are available (Pharmacia Biotech, Inc., Piscataway, NJ). In an alternative, a polypeptide/Fc polypeptide (as discussed above) is attached to Protein A- or Protein Gcontaining chromatography columns through interaction with the Fc moiety. The HPR1 and HPR2 polypeptides also find use in identifying cells that express a binding partner on the cell surface. Polypeptides are bound to a solid phase such as a column chromatography matrix or a similar suitable substrate. For example, magnetic microspheres can be coated with the polypeptides and held in an incubation vessel through a magnetic field. Suspensions of cell mixtures containing potential bindingpartner-expressing cells are contacted with the solid phase having the polypeptides thereon. Cells expressing the binding partner on the cell surface bind to the fixed polypeptides, and unbound cells are washed away. Alternatively, HPR1 and HPR2 polypeptides can be conjugated to a detectable moiety, then incubated with cells to be tested for binding partner expression. After incubation, unbound labeled matter is removed and the presence or absence of the detectable moiety on the cells is determined. In a further alternative, mixtures of cells suspected of expressing the binding partner are incubated with biotinylated polypeptides. Incubation periods are typically at least one hour in duration to ensure sufficient binding. The resulting mixture then is passed through a column packed with avidin-coated beads, whereby the high affinity of biotin for avidin provides binding of the desired cells to the beads. Procedures for using avidin-coated beads are known (see Berenson, et al. J. Cell. Biochem., 10D:239, 1986). Washing to remove unbound material, and the release of the bound cells,

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are performed using conventional methods. In some instances, the above methods for screening for or identifying binding partners may also be used or modified to isolate or purify such binding partner molecules or cells expressing them.

Measuring Biological Activity. HPR1 and HPR2 polypeptides also find use in measuring the biological activity of HPR1-binding and/or HPR2-binding polypeptides in terms of their binding affinity. The polypeptides thus can be employed by those conducting "quality assurance" studies, e.g., to monitor shelf life and stability of polypeptide under different conditions. For example, the polypeptides can be employed in a binding affinity study to measure the biological activity of a binding partner polypeptide that has been stored at different temperatures, or produced in different cell types. The polypeptides also can be used to determine whether biological activity is retained after modification of a binding partner polypeptide (e.g., chemical modification, truncation, mutation, etc.). The binding affinity of the modified polypeptide is compared to that of an unmodified binding polypeptide to detect any adverse impact of the modifications on biological activity of the binding polypeptide. The biological activity of a binding polypeptide thus can be ascertained before it is used in a research study, for example.

Carriers and Delivery Agents. The polypeptides also find use as carriers for delivering agents attached thereto to cells bearing identified binding partners. The polypeptides thus can be used to deliver diagnostic or therapeutic agents to such cells (or to other cell types found to express binding partners on the cell surface) in in vitro or in vivo procedures. Detectable (diagnostic) and therapeutic agents that can be attached to a polypeptide include, but are not limited to, toxins, other cytotoxic agents, drugs, radionuclides, chromophores, enzymes that catalyze a colorimetric or fluorometric reaction, and the like, with the particular agent being chosen according to the intended application. Among the toxins are ricin, abrin, diphtheria toxin, Pseudomonas aeruginosa exotoxin A, ribosomal inactivating polypeptides, mycotoxins such as trichothecenes, and derivatives and fragments (e.g., single chains) thereof. Radionuclides suitable for diagnostic use include, but are not limited to, <sup>123</sup>I, <sup>131</sup>I, <sup>99m</sup>Tc, <sup>111</sup>In, and <sup>76</sup>Br. Examples of radionuclides suitable for therapeutic use are <sup>131</sup>I, <sup>211</sup>At, <sup>77</sup>Br, <sup>186</sup>Re, <sup>188</sup>Re, <sup>212</sup>Pb, <sup>212</sup>Bi, <sup>109</sup>Pd, <sup>64</sup>Cu, and <sup>67</sup>Cu. Such agents can be attached to the polypeptide by any suitable conventional procedure. The polypeptide comprises functional groups on amino acid side chains that can be reacted with functional groups on a desired agent to form covalent bonds, for example. Alternatively, the polypeptide or agent can be derivatized to generate or attach a desired reactive functional group. The derivatization can involve attachment of one of the bifunctional coupling reagents available for attaching various molecules to polypeptides (Pierce Chemical Company, Rockford, Illinois). A number of techniques for radiolabeling polypeptides are known. Radionuclide metals can be attached to polypeptides by using a suitable bifunctional chelating agent, for example. Conjugates comprising polypeptides and a suitable diagnostic or therapeutic agent (preferably covalently linked) are thus prepared. The conjugates are administered or otherwise employed in an amount appropriate for the particular application.

### 5 Treating Diseases Using HPR1 and/or HPR2 Polypeptides and Antagonists Thereof

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It is anticipated that the HPR1 and HPR2 polypeptides, fragments, variants, antagonists, agonists, antibodies, and binding partners of the invention will be useful for treating medical conditions and diseases including, but not limited to, cell proliferation, metabolic, and reproductive hormone related conditions as described further herein. The therapeutic molecule or molecules to be used will depend on the etiology of the condition to be treated and the biological pathways involved, and variants, fragments, and binding partners of HPR1 and/or HPR2 polypeptides may have effects similar to or different from HPR1 or HPR2 polypeptides. For example, an antagonist of the ligand-binding activity of HPR1 and/or HPR2 polypeptides may be selected for treatment of conditions involving ligand-binding activity, but a particular fragment of a given HPR1 or HPR2 polypeptide may also act as an effective dominant negative antagonist of that activity. Therefore, in the following paragraphs "HPR1 and HPR2 polypeptides or antagonists" refers to all HPR1 and HPR2 polypeptides, fragments, variants, antagonists, agonists, antibodies, and binding partners etc. of the invention, and it is understood that a specific molecule or molecules can be selected from those provided as embodiments of the invention by individuals of skill in the art, according to the biological and therapeutic considerations described herein.

Provided herein are methods for using HPR1 and HPR2 polypeptides or antagonists, compositions or combination therapies to treat various hematologic and oncologic disorders. For example, HPR1 and HPR2 polypeptides or antagonists are used to treat various forms of cancer, including acute myelogenous leukemia, Epstein-Barr virus-positive nasopharyngeal carcinoma, glioma, colon, stomach, prostate, renal cell, cervical and ovarian cancers, lung cancer (SCLC and NSCLC), including cancer-associated cachexia, fatigue, asthenia, paraneoplastic syndrome of cachexia and hypercalcemia. Additional diseases treatable with the subject HPR1 and HPR2 polypeptides or antagonists, compositions or combination therapies are solid tumors, including sarcoma, osteosarcoma, and carcinoma, such as adenocarcinoma (for example, breast cancer) and squamous cell carcinoma. In addition, the subject compounds, compositions or combination therapies are useful for treating leukemia, including acute myelogenous leukemia, chronic or acute lymphoblastic leukemia and hairy cell leukemia. Other malignancies with invasive metastatic potential can be treated with the subject compounds, compositions and combination therapies, including multiple myeloma. In addition, the disclosed HPR1 and HPR2 polypeptides or antagonists, compositions and combination therapies can be used to treat anemias and hematologic disorders, including anemia of chronic disease, aplastic anemia, including Fanconi's aplastic anemia; idiopathic thrombocytopenic purpura (ITP); myelodysplastic syndromes (including refractory anemia, refractory anemia with ringed sideroblasts, refractory anemia with excess blasts, refractory anemia with excess blasts in transformation); myelofibrosis/myeloid metaplasia; and sickle cell vasocclusive crisis.

Various lymphoproliferative disorders also are treatable with the disclosed HPR1 and HPR2 polypeptides or antagonists, compositions or combination therapies. These include, but are not limited to autoimmune lymphoproliferative syndrome (ALPS), chronic lymphoblastic leukemia, hairy cell

leukemia, chronic lymphatic leukemia, peripheral T-cell lymphoma, small lymphocytic lymphoma, mantle cell lymphoma, follicular lymphoma, Burkitt's lymphoma, Epstein-Barr virus-positive T cell lymphoma, histiocytic lymphoma, Hodgkin's disease, diffuse aggressive lymphoma, acute lymphatic leukemias, T gamma lymphoproliferative disease, cutaneous B cell lymphoma, cutaneous T cell lymphoma (i.e., mycosis fungoides) and Sézary syndrome.

In addition, the subject invention provides HPR1 and HPR2 polypeptides or antagonists, compositions and combination therapies for the treatment of non-arthritic medical conditions of the bones and joints. This encompasses osteoclast disorders that lead to bone loss, such as but not limited to osteoporosis, including post-menopausal osteoporosis, periodontitis resulting in tooth loosening or loss, and prosthesis loosening after joint replacement (generally associated with an inflammatory response to wear debris). This latter condition also is called "orthopedic implant osteolysis." Another condition treatable by administering HPR1 and HPR2 polypeptides or antagonists, is temporal mandibular joint dysfunction (TMJ).

The disclosed HPR1 and HPR2 polypeptides or antagonists, compositions and combination therapies furthermore are useful for treating neurodegenerative conditions such as acute polyneuropathy; anorexia nervosa; Bell's palsy; chronic fatigue syndrome; transmissible dementia, including Creutzfeld-Jacob disease; demyelinating neuropathy; Guillain-Barre syndrome; vertebral disc disease; Gulf war syndrome; myasthenia gravis; chronic neuronal degeneration; and stroke, including cerebral ischemic diseases.

# 25 Administration of HPR1 and HPR2 Polypeptides and Antagonists Thereof

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This invention provides compounds, compositions, and methods for treating a patient, preferably a mammalian patient, and most preferably a human patient, who is suffering from a medical disorder, and in particular an HPR1- or HPR2-mediated disorder. Such HPR1- or HPR2-mediated disorders include conditions caused (directly or indirectly) or exacerbated by binding between HPR1 and/or HPR2 and a binding partner. For purposes of this disclosure, the terms "illness," "disease," "medical condition," "abnormal condition" and the like are used interchangeably with the term "medical disorder." The terms "treat", "treating", and "treatment" used herein includes curative, preventative (e.g., prophylactic) and palliative or ameliorative treatment. For such therapeutic uses, HPR1 and HPR2 polypeptides and fragments, HPR1 and HPR2 nucleic acids encoding the HPR1 and HPR2 polypeptides, and/or agonists or antagonists of the HPR1 and/or HPR2 polypeptides such as antibodies can be administered to the patient in need through well-known means. Compositions of the present invention can contain a polypeptide in any form described herein, such as native polypeptides, variants, derivatives, oligomers, and biologically active fragments. In particular embodiments, the composition comprises a soluble polypeptide or an oligomer comprising soluble HPR1 and/or HPR2 polypeptides.

<u>Therapeutically Effective Amount</u>. In practicing the method of treatment or use of the present invention, a therapeutically effective amount of a therapeutic agent of the present invention is

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administered to a patient having a condition to be treated, preferably to treat or ameliorate diseases associated with the activity of an HPR1 and/or HPR2 polypeptide. "Therapeutic agent" includes without limitation any of the HPR1 or HPR2 polypeptides, fragments, and variants; nucleic acids encoding the HPR1 and HPR2 polypeptides, fragments, and variants; agonists or antagonists of the HPR1 and HPR2 polypeptides such as antibodies; HPR1 and/or HPR2 polypeptide binding partners; complexes formed from the HPR1 and/or HPR2 polypeptides, fragments, variants, and binding partners, etc. As used herein, the term "therapeutically effective amount" means the total amount of each therapeutic agent or other active component of the pharmaceutical composition or method that is sufficient to show a meaningful patient benefit, i.e., treatment, healing, prevention or amelioration of the relevant medical condition, or an increase in rate of treatment, healing, prevention or amelioration of such conditions. When applied to an individual therapeutic agent or active ingredient, administered alone, the term refers to that ingredient alone. When applied to a combination, the term refers to combined amounts of the ingredients that result in the therapeutic effect, whether administered in combination, serially or simultaneously. As used herein, the phrase "administering a therapeutically effective amount" of a therapeutic agent means that the patient is treated with said therapeutic agent in an amount and for a time sufficient to induce an improvement, and preferably a sustained improvement, in at least one indicator that reflects the severity of the disorder. An improvement is considered "sustained" if the patient exhibits the improvement on at least two occasions separated by one or more weeks. The degree of improvement is determined based on signs or symptoms, and determinations can also employ questionnaires that are administered to the patient, such as quality-oflife questionnaires. Various indicators that reflect the extent of the patient's illness can be assessed for determining whether the amount and time of the treatment is sufficient. The baseline value for the chosen indicator or indicators is established by examination of the patient prior to administration of the first dose of the therapeutic agent. Preferably, the baseline examination is done within about 60 days of administering the first dose. If the therapeutic agent is being administered to treat acute symptoms, the first dose is administered as soon as practically possible after the injury has occurred. Improvement is induced by administering therapeutic agents such as HPR1 and/or HPR2 polypeptides or antagonists until the patient manifests an improvement over baseline for the chosen indicator or indicators. In treating chronic conditions, this degree of improvement is obtained by repeatedly administering this medicament over a period of at least a month or more, e.g., for one, two, or three months or longer, or indefinitely. A period of one to six weeks, or even a single dose, often is sufficient for treating injuries or acute conditions. Although the extent of the patient's illness after treatment may appear improved according to one or more indicators, treatment may be continued indefinitely at the same level or at a reduced dose or frequency. Once treatment has been reduced or discontinued, it later may be resumed at the original level if symptoms should reappear.

<u>Dosing.</u> One skilled in the pertinent art will recognize that suitable dosages will vary, depending upon such factors as the nature and severity of the disorder to be treated, the patient's body weight, age, general condition, and prior illnesses and/or treatments, and the route of administration.

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Preliminary doses can be determined according to animal tests, and the scaling of dosages for human administration is performed according to art-accepted practices such as standard dosing trials. For example, the therapeutically effective dose can be estimated initially from cell culture assays. The dosage will depend on the specific activity of the compound and can be readily determined by routine experimentation. A dose can be formulated in animal models to achieve a circulating plasma concentration range that includes the IC50 (i.e., the concentration of the test compound which achieves a half-maximal inhibition of symptoms) as determined in cell culture, while minimizing toxicities. Such information can be used to more accurately determine useful doses in humans. Ultimately, the attending physician will decide the amount of polypeptide of the present invention with which to treat each individual patient. Initially, the attending physician will administer low doses of polypeptide of the present invention and observe the patient's response. Larger doses of polypeptide of the present invention can be administered until the optimal therapeutic effect is obtained for the patient, and at that point the dosage is not increased further. It is contemplated that the various pharmaceutical compositions used to practice the method of the present invention should contain about 0.01 ng to about 100 mg (preferably about 0.1 ng to about 10 mg, more preferably about 0.1 microgram to about 1 mg) of polypeptide of the present invention per kg body weight. In one embodiment of the invention, HPR1 and/or HPR2 polypeptides or antagonists are administered one time per week to treat the various medical disorders disclosed herein, in another embodiment is administered at least two times per week, and in another embodiment is administered at least three times per week. If injected, the effective amount of HPR1 or HPR2 polypeptides or antagonists per adult dose ranges from 1-20 mg/m<sup>2</sup>, and preferably is about 5-12 mg/m<sup>2</sup>. Alternatively, a flat dose can be administered, whose amount may range from 5-100 mg/dose. Exemplary dose ranges for a flat dose to be administered by subcutaneous injection are 5-25 mg/dose, 25-50 mg/dose and 50-100 mg/dose. In one embodiment of the invention, the various indications described below are treated by administering a preparation acceptable for injection containing HPR1 and/or HPR2 polypeptides or antagonists at 25 mg/dose, or alternatively, containing 50 mg per dose. The 25 mg or 50 mg dose can be administered repeatedly, particularly for chronic conditions. If a route of administration other than injection is used, the dose is appropriately adjusted in accord with standard medical practices. In many instances, an improvement in a patient's condition will be obtained by injecting a dose of about 25 mg of HPR1 or HPR2 polypeptides or antagonists one to three times per week over a period of at least three weeks, or a dose of 50 mg of HPR1 or HPR2 polypeptides or antagonists one or two times per week for at least three weeks, though treatment for longer periods may be necessary to induce the desired degree of improvement. For incurable chronic conditions, the regimen can be continued indefinitely, with adjustments being made to dose and frequency if such are deemed necessary by the patient's physician. The foregoing doses are examples for an adult patient who is a person who is 18 years of age or older. For pediatric patients (age 4-17), a suitable regimen involves the subcutaneous injection of 0.4 mg/kg, up to a maximum dose of 25 mg of HPR1 or HPR2 polypeptides or antagonists, administered by subcutaneous injection one or more times per week. If an antibody against an HPR1 and/or HPR2 polypeptide is used as the HPR1

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and/or HPR2 polypeptide antagonist, a preferred dose range is 0.1 to 20 mg/kg, and more preferably is 1-10 mg/kg. Another preferred dose range for an anti-HPR1 polypeptide and/or anti-HPR2 polypeptide antibody is 0.75 to 7.5 mg/kg of body weight. Humanized antibodies are preferred, that is, antibodies in which only the antigen-binding portion of the antibody molecule is derived from a non-human source. Such antibodies can be injected or administered intravenously.

Formulations. Compositions comprising an effective amount of an HPR1 and/or HPR2 polypeptide of the present invention (from whatever source derived, including without limitation from recombinant and non-recombinant sources), in combination with other components such as a physiologically acceptable diluent, carrier, or excipient, are provided herein. The term "pharmaceutically acceptable" means a non-toxic material that does not interfere with the effectiveness of the biological activity of the active ingredient(s). Formulations suitable for administration include aqueous and non-aqueous sterile injection solutions which can contain anti-oxidants, buffers, bacteriostats and solutes which render the formulation isotonic with the blood of the recipient; and aqueous and non-aqueous sterile suspensions which can include suspending agents or thickening The polypeptides can be formulated according to known methods used to prepare pharmaceutically useful compositions. They can be combined in admixture, either as the sole active material or with other known active materials suitable for a given indication, with pharmaceutically acceptable diluents, preservatives, emulsifiers, solubilizers, adjuvants and/or carriers. formulations for pharmaceutical compositions include those described in Remington's Pharmaceutical Sciences, 16th ed. 1980, Mack Publishing Company, Easton, PA. In addition, such compositions can be complexed with polyethylene glycol (PEG), metal ions, or incorporated into polymeric compounds such as polyacetic acid, polyglycolic acid, hydrogels, dextran, etc., or incorporated into liposomes, microemulsions, micelles, unilamellar or multilamellar vesicles, erythrocyte ghosts or spheroblasts. Suitable lipids for liposomal formulation include, without limitation, monoglycerides, diglycerides, sulfatides, lysolecithin, phospholipids, saponin, bile acids, and the like. Preparation of such liposomal formulations is within the level of skill in the art, as disclosed, for example, in U.S. Pat. No. 4,235,871; U.S. Pat. No. 4,501,728; U.S. Pat. No. 4,837,028; and U.S. Pat. No. 4,737,323. Such compositions will influence the physical state, solubility, stability, rate of in vivo release, and rate of in vivo clearance, and are thus chosen according to the intended application, so that the characteristics of the carrier will depend on the selected route of administration. In one preferred embodiment of the invention, sustained-release forms of HPR1 and/or HPR2 polypeptides are used. Sustained-release forms suitable for use in the disclosed methods include, but are not limited to, HPR1 and/or HPR2 polypeptides that are encapsulated in a slowly-dissolving biocompatible polymer (such as the alginate microparticles described in U.S. No. 6,036,978), admixed with such a polymer (including topically applied hydrogels), and or encased in a biocompatible semi-permeable implant.

<u>Combinations of Therapeutic Compounds.</u> An HPR1 or HPR2 polypeptide of the present invention may be active in multimers (e.g., heterodimers or homodimers) or complexes with itself or other polypeptides. As a result, pharmaceutical compositions of the invention may comprise a

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polypeptide of the invention in such multimeric or complexed form. The pharmaceutical composition of the invention may be in the form of a complex of the polypeptide(s) of present invention along with polypeptide or peptide antigens. The invention further includes the administration of HPR1 and/or HPR2 polypeptides or antagonists concurrently with one or more other drugs that are administered to the same patient in combination with the HPR1 and/or HPR2 polypeptides or antagonists, each drug being administered according to a regimen suitable for that medicament. "Concurrent administration" encompasses simultaneous or sequential treatment with the components of the combination, as well as regimens in which the drugs are alternated, or wherein one component is administered long-term and the other(s) are administered intermittently. Components can be administered in the same or in separate compositions, and by the same or different routes of administration. Examples of components that can be included in the pharmaceutical composition of the invention are: cytokines, lymphokines, or other hematopoietic factors such as M-CSF, GM-CSF, TNF, IL-1, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-11, IL-12, IL-13, IL-14, IL-15, IL-17, IL-18, IL-23, IFN, TNF0, TNF1, TNF2, G-CSF, Meg-CSF, thrombopoietin, stem cell factor, and erythropoietin. The pharmaceutical composition can further contain other agents which either enhance the activity of the polypeptide or compliment its activity or use in treatment. Such additional factors and/or agents may be included in the pharmaceutical composition to produce a synergistic effect with polypeptide of the invention, or to minimize side effects. Additional examples of drugs to be administered concurrently include but are not limited to antivirals, antibiotics, analgesics, corticosteroids, antagonists of inflammatory cytokines, non-steroidal anti-inflammatories, pentoxifylline, thalidomide, and disease-modifying antirheumatic drugs (DMARDs) such as azathioprine, cyclophosphamide, cyclosporine, hydroxychloroquine sulfate, methotrexate, leflunomide, minocycline, penicillamine, sulfasalazine and gold compounds such as oral gold, gold sodium thiomalate, and aurothioglucose. Additionally, HPR1 and/or HPR2 polypeptides or antagonists can be combined with a second HPR1 and/or HPR2 polypeptide/antagonist, including an antibody against an HPR1 and/or HPR2 polypeptide, or an HPR1 polypeptide-derived peptide or HPR2 polypeptide-derived peptide that acts as a competitive inhibitor of native HPR1 and/or HPR2 polypeptides.

Routes of Administration. Any efficacious route of administration may be used to therapeutically administer HPR1 and HPR2 polypeptides or antagonists thereof, including those compositions comprising nucleic acids. Parenteral administration includes injection, for example, via intra-articular, intravenous, intramuscular, intralesional, intraperitoneal or subcutaneous routes by bolus injection or by continuous infusion., and also includes localized administration, e.g., at a site of disease or injury. Other suitable means of administration include sustained release from implants; aerosol inhalation and/or insufflation.; eyedrops; vaginal or rectal suppositories; buccal preparations; oral preparations, including pills, syrups, lozenges or chewing gum; and topical preparations such as lotions, gels, sprays, ointments or other suitable techniques. Alternatively, polypeptideaceous HPR1 and HPR2 polypeptides or antagonists may be administered by implanting cultured cells that express the polypeptide, for example, by implanting cells that express HPR1 and/or HPR2 polypeptides or

antagonists. Cells may also be cultured ex vivo in the presence of polypeptides of the present invention in order to proliferate or to produce a desired effect on or activity in such cells. Treated cells can then be introduced in vivo for therapeutic purposes. In another embodiment, the patient's own cells are induced to produce HPR1 and/or HPR2 polypeptides or antagonists by transfection *in vivo* or *ex vivo* with a DNA that encodes HPR1 and/or HPR2 polypeptides or antagonists. This DNA can be introduced into the patient's cells, for example, by injecting naked DNA or liposome-encapsulated DNA that encodes HPR1 and/or HPR2 polypeptides or antagonists, or by other means of transfection. Nucleic acids of the invention can also be administered to patients by other known methods for introduction of nucleic acid into a cell or organism (including, without limitation, in the form of viral vectors or naked DNA). When HPR1 and/or HPR2 polypeptides or antagonists are administered in combination with one or more other biologically active compounds, these can be administered by the same or by different routes, and can be administered simultaneously, separately or sequentially.

Oral Administration. When a therapeutically effective amount of polypeptide of the present invention is administered orally, polypeptide of the present invention will be in the form of a tablet, capsule, powder, solution or elixir. When administered in tablet form, the pharmaceutical composition of the invention can additionally contain a solid carrier such as a gelatin or an adjuvant. The tablet, capsule, and powder contain from about 5 to 95% polypeptide of the present invention, and preferably from about 25 to 90% polypeptide of the present invention. When administered in liquid form, a liquid carrier such as water, petroleum, oils of animal or plant origin such as peanut oil, mineral oil, soybean oil, or sesame oil, or synthetic oils can be added. The liquid form of the pharmaceutical composition can further contain physiological saline solution, dextrose or other saccharide solution, or glycols such as ethylene glycol, propylene glycol or polyethylene glycol. When administered in liquid form, the pharmaceutical composition contains from about 0.5 to 90% by weight of polypeptide of the present invention, and preferably from about 1 to 50% polypeptide of the present invention.

Intravenous Administration. When a therapeutically effective amount of polypeptide of the present invention is administered by intravenous, cutaneous or subcutaneous injection, polypeptide of the present invention will be in the form of a pyrogen-free, parenterally acceptable aqueous solution. The preparation of such parenterally acceptable polypeptide solutions, having due regard to pH, isotonicity, stability, and the like, is within the skill in the art. A preferred pharmaceutical composition for intravenous, cutaneous, or subcutaneous injection should contain, in addition to polypeptide of the present invention, an isotonic vehicle such as Sodium Chloride Injection, Ringer's Injection, Dextrose Injection, Dextrose and Sodium Chloride Injection, Lactated Ringer's Injection, or other vehicle as known in the art. The pharmaceutical composition of the present invention can also contain stabilizers, preservatives, buffers, antioxidants, or other additives known to those of skill in the art. The duration of intravenous therapy using the pharmaceutical composition of the present invention will vary, depending on the severity of the disease being treated and the condition and potential idiosyncratic response of each individual patient. It is contemplated that the duration of each application of the polypeptide of the present invention will be in the range of 12 to 24 hours of continuous intravenous

administration. Ultimately the attending physician will decide on the appropriate duration of intravenous therapy using the pharmaceutical composition of the present invention.

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Bone and Tissue Administration. For compositions of the present invention which are useful for bone, cartilage, tendon or ligament disorders, the therapeutic method includes administering the composition topically, systematically, or locally as an implant or device. When administered, the therapeutic composition for use in this invention is, of course, in a pyrogen-free, physiologically acceptable form. Further, the composition can desirably be encapsulated or injected in a viscous form for delivery to the site of bone, cartilage or tissue damage. Topical administration may be suitable for wound healing and tissue repair. Therapeutically useful agents other than a polypeptide of the invention which can also optionally be included in the composition as described above, can alternatively or additionally, be administered simultaneously or sequentially with the composition in the methods of the invention. Preferably for bone and/or cartilage formation, the composition would include a matrix capable of delivering the polypeptide-containing composition to the site of bone and/or cartilage damage, providing a structure for the developing bone and cartilage and optimally capable of being resorbed into the body. Such matrices can be formed of materials presently in use for other implanted medical applications. The choice of matrix material is based on biocompatibility, biodegradability, mechanical properties, cosmetic appearance and interface properties. The particular application of the compositions will define the appropriate formulation. Potential matrices for the compositions can be biodegradable and chemically defined calcium sulfate, tricalciumphosphate, hydroxyapatite, polylactic acid, polyglycolic acid and polyanhydrides. Other potential materials are biodegradable and biologically well-defined, such as bone or dermal collagen. Further matrices are comprised of pure polypeptides or extracellular matrix components. Other potential matrices are nonbiodegradable and chemically defined, such as sintered hydroxapatite, bioglass, aluminates, or other ceramics Matrices can be comprised of combinations of any of the above mentioned types of material, such as polylactic acid and hydroxyapatite or collagen and tricalciumphosphate. Presently preferred is a 50:50 (mole weight) copolymer of lactic acid and glycolic acid in the form of porous particles having diameters ranging from 150 to 800 microns. In some applications, it will be useful to utilize a sequestering agent, such as carboxymethyl cellulose or autologous blood clot, to prevent the polypeptide compositions from disassociating from the matrix. A preferred family of sequestering agents is cellulosic materials such as alkylcelluloses, the most preferred being cationic salts of carboxymethylcellulose (CMC). Other preferred sequestering agents include hyaluronic acid, sodium alginate, poly(ethylene glycol), polyoxyethylene oxide, carboxyvinyl polymer and poly(vinyl alcohol). The amount of sequestering agent useful herein is 0.5-20 wt %, preferably 1-10 wt % based on total formulation weight, which represents the amount necessary to prevent desorbtion of the polypeptide from the polymer matrix and to provide appropriate handling of the composition, yet not so much that the progenitor cells are prevented from infiltrating the matrix, thereby providing the polypeptide the opportunity to assist the osteogenic activity of the progenitor cells. In further compositions, polypeptides of the invention can be combined with other agents beneficial to the treatment of the bone

and/or cartilage defect, wound, or tissue in question. Progress can be monitored by periodic assessment of tissue/bone growth and/or repair, for example, X-rays, histomorphometric determinations and tetracycline labeling.

<u>Veterinary Uses.</u> In addition to human patients, HPR1 and HPR2 polypeptides and antagonists are useful in the treatment of disease conditions in non-human animals, such as pets (dogs, cats, birds, primates, etc.), domestic farm animals (horses cattle, sheep, pigs, birds, etc.), or any animal that suffers from a TNFα-mediated inflammatory or arthritic condition. In such instances, an appropriate dose can be determined according to the animal's body weight. For example, a dose of 0.2-1 mg/kg may be used. Alternatively, the dose is determined according to the animal's surface area, an exemplary dose ranging from 0.1-20 mg/m², or more preferably, from 5-12 mg/m². For small animals, such as dogs or cats, a suitable dose is 0.4 mg/kg. In a preferred embodiment, HPR1 and/or HPR2 polypeptides or antagonists (preferably constructed from genes derived from the same species as the patient), are administered by injection or other suitable route one or more times per week until the animal's condition is improved, or they can be administered indefinitely.

Manufacture of Medicaments. The present invention also relates to the use of HPR1 and HPR2 polypeptides, fragments, and variants; nucleic acids encoding the HPR1 or HPR2 polypeptides, fragments, and variants; agonists or antagonists of the HPR1 and/or HPR2 polypeptides such as antibodies; HPR1 and/or HPR2 polypeptide binding partners; complexes formed from the HPR1 and/or HPR2 polypeptides, fragments, variants, and binding partners, etc, in the manufacture of a medicament for the prevention or therapeutic treatment of each medical disorder disclosed herein.

25 <u>EXAMPLES</u>

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The following examples are intended to illustrate particular embodiments and not to limit the scope of the invention.

#### **EXAMPLE 1**

# A. Identification of HPR1, a New Member of the Human Hematopoietin Receptor Family

A data set was received from Celera Genomics (Rockville, Maryland) containing a listing of amino acid sequences predicted to be encoded by the human genome. This data set was searched with a BLAST algorithm to identify hematopoietin receptor family polypeptides. Several amino acid sequences, including two overlapping amino acid sequences (SEQ ID NO: 1 and SEQ ID NO:2), were identified as comprising partial amino acid sequences of a new human hematopoietin receptor polypeptide, HPR1. These amino acids sequences were used to identify a DNA sequence (SEQ ID NO:3) encoding an HPR1 polypeptide having the amino acid sequence shown in SEQ ID NO:4; nucleotides 132 through 2366 of SEQ ID NO:3 encode SEQ ID NO:4, with nucleotides 2367 through 2369 corresponding to a stop codon. The HPR1 coding sequence (nucleotides 132 through 2369 of SEQ ID NO:3) is presented as SEQ ID NO:5. The HPR1 sequences of SEQ ID NOs 3 and 5 were confirmed by three independent PCR amplification experiments from a U937 cDNA library. These HPR1 coding sequences were compared with publicly available preliminary human genomic DNA sequences, and the following chromosome 5 contigs were identified as containing HPR1 coding sequences: ACO22265.3, ACO08914.3, ACO08857.4, and ACO16596.4. The human genomic region

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corresponding to these contigs also includes the gene for gp130, which suggests that gp130 and HPR1 may derive from a common ancestral gene by gene duplication. The approximate positions of the exons containing HPR1 coding sequence in the AC022265.3 contig are shown in the table below, along with their locations relative to SEQ ID NOs 3 and 5; note that the 5' and 3' untranslated regions may extend further along the contig sequence beyond those portions that correspond to SEQ ID NOs 3 and 5, as indicated by the parentheses around the AC022265.3 endpoints in the table. Due to the preliminary sequence and assembly of the contig sequence, the exons within the contig are not always in the right order or orientation with respect to each other, and may contain sequence variations due to inaccurate sequence data or allelic polymorphism.

Corresponding positions of HPR1 gene exons in human contig AC022265.3 and in cDNA sequences:

	Position in AC022265.3	Position in SEQ ID NO:3 / Position in SEQ ID NO:5
Exon I	(128423)-128559	1-137 / 1-6
Exon 2	134501-134591	138-228 / 7-97
Exon 3	143777-143894	229-346 / 98-215
Exon 4	147256-147437	347-528 / 216-397
Exon 5	51249-51098	529-680 / 398-549
Exon 6	44322-44157	681-846 / 550-715
Exon 7	16473-16394	847-926 / 716-795
Exon 8	30331-30115	927-1143 / 796-1012
Exon 9	178626-178808	1144-1326 / 1013-1195
Exon 10	179879-179980	1327-1428 / 1196-1297
Exon 11	180785-180931	1429-1575 / 1298-1444
Exon 12	183052-183192	1576-1716 / 1445-1585
Exon 13	185997-186090	1717-1810 / 1586-1679
Exon 14	187367-187448	1811-1892 / 1680-1761
Exon 15	189165-(189747)	1893-2480 / 1762-2238

A nucleic acid encoding a polypeptide with a high degree of amino acid similarity (approximately 61% amino acid identity) to human HPR1 was isolated from Mus musculus. The Mus HPR1 amino acid sequence is presented as SEQ ID NO:12, and due to its high level of similarity with human HPR1, is considered to be the murine homologue of human HPR1. PCR amplification of cDNA sequences corresponding to mRNAs encoding murine HPR1 identified a cDNA molecule encoding SEQ ID NO:12; the nucleotide sequence of this murine HPR1 cDNA is presented as SEQ ID NO:28. Nucleotides 1 through 2178 of SEQ ID NO:28 encode SEQ ID NO:12, with nucleotides 2179-2181 corresponding to a stop codon. Variants of the murine HPR1 amino acid sequence that are likely allelic variants have been identified in which the 'T' residue at position 121 of SEQ ID NO:28 is changed to a 'C' residue, resulting in a change from the Phe residue at position 41 of SEQ ID NO:4 to a Leu residue, and in which the 'G' residue at position 1666 of SEQ ID NO:28 is changed to an 'A' residue, resulting in a change from the Asp residue at position 556 of SEQ ID NO:4 to an Asn residue.

Several splice variations of the HPR1 sequences have been identified in human genomic sequences and are included within the scope of the invention. For example, amino acids 1 through 55 of SEQ ID NO:1 match the amino acid sequence of HPR1 presented in SEQ ID NO:4, while amino

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acids 56 through 77 of SEQ ID NO:1 may be a portion of an alternatively spliced exon added following the exon/intron boundary identified between nucleotides 846 and 847 of SEQ ID NO:3 (nucleotides 715 and 716 of SEQ ID NO:5). In an additional potential splice variant, an amino acid sequence ending in the amino acids of SEQ ID NO:10 could be substituted for the amino acids leading up to and including the lysine at position 190 of SEQ ID NO:4. However, such a splice variant would require an additional exon/intron boundary approximately between nucleotides 701 and 702 of SEQ ID NO:3 (nucleotides 570 and 571 of SEQ ID NO:5). In a further potential splice variant, the amino acid sequence of SEQ ID NO:11 could be substituted for amino acids 238 through 266 of SEQ ID NO:4 by replacing exon 7 with an alternative exon encoding the SEQ ID NO:11 amino acids. In this potential variant, 29 amino acids C-terminal to the WSXWS motif and including the N-terminal portion of the most N-terminal fibronectin type III repeat (as shown in Table 1) would be replaced with 15 amino acids, resulting in deletion of a portion of the most N-terminal fibronectin type III repeat, including two highly conserved Trp residues.

Additional variations of HPR1 polypeptides are provided as naturally occurring genomic variants of the HPR1 sequences disclosed herein; such variations may be incorporated into an HPR1 polypeptide or nucleic acid individually or in any combination, or in combination with alternative splice variation as described above. As one example, amino acids 5 through 40 of SEQ ID NO:2 match SEQ ID NO:4, with amino acid 4 of SEQ ID NO:2 likely representing an allelic variation, where the change from the Asn residue position 187 of SEQ ID NO:4 to a Thr residue in SEQ ID NO:2 could be caused by a single change from 'A' to 'C' at position 691 of SEQ ID NO:3 or 560 of SEQ ID NO:5. This variation and others are listed in the table below:

Amino Acid Change	Position in SEQ ID NO:4	Nucleotide Change	Position in SEQ ID NO:3 / Position in SEQ ID NO:5
Thr -> Ala	83	A -> G	378 / 247
Asp -> Asn	168	G -> A	633 / 502
Asn -> Thr	187	A -> C	691 / 560
Ser -> Pro	361	T -> C	1212 / 1081
Ala -> Gly	362	C -> G	1216 / 1085
Ser -> Asn	510	G -> A	1660 / 1529
Asn -> Asp	517	A -> G	1680 / 1549
Arg -> Gly	679	A -> G	2166 / 2035

## B. Identification of HPR2, a New Member of the Human Hematopoietin Receptor Family

A data set was received from Celera Genomics (Rockville, Maryland) containing a listing of amino acid sequences predicted to be encoded by the human genome. This data set was searched with a BLAST algorithm to identify hematopoietin receptor family polypeptides. Several amino acid sequences, including SEQ ID NO:16, were identified as comprising partial amino acid sequences of a new human hematopoietin receptor polypeptide, HPR2. These amino acids sequences were used to identify a DNA sequence (SEQ ID NO:19) encoding an HPR2 polypeptide having the amino acid sequence shown in SEQ ID NO:21; nucleotides 107 through 1993 of SEQ ID 19 encode SEQ ID

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NO:21, with nucleotides 1994 through 1996 corresponding to a stop codon. The HPR2 coding sequence (nucleotides 107 through 1996 of SEQ ID NO:19) is presented as SEQ ID NO:20. The HPR2 sequences of SEQ ID NOs 19 and 20 were confirmed by independent PCR amplification experiments from a human lymph node cDNA library and a CB23 B cell line cDNA library. These PCR amplification experiments also identified two additional splice variants of the HPR2 cDNA sequence referred to as HPR2-ex8-ex9 and HPR2-ex9; the coding sequences for HPR2-ex8-ex9 and HPR2-ex9 are presented as SEQ ID NOs 22 and 24, respectively, and the amino acid sequences they encode are presented as SEQ ID NOs 23 and 25, respectively. The HPR2 cDNA sequences of SEQ ID NOs 19, 20, and the HPR2-ex8-ex9 cDNA of SEQ ID NO:22 were present in both the lymph node and CB23 cDNA libraries, while the HPR2-ex9 cDNA of SEQ ID NO:24 was only present in the lymph node library.

These HPR2 coding sequences were compared with publicly available preliminary human genomic DNA sequences, and the following chromosome 1 contigs were identified as containing HPR2 coding sequences: GenBank accession numbers AL109843 (1p31.2-32.1) and AL389925. The human genomic region corresponding to the AL389925 contig also includes the gene for IL-12RB2, which suggests that IL-12RB2 and HPR2 may derive from a common ancestral gene by gene duplication. The approximate positions of the exons containing HPR2 coding sequence in the AL109843 and AL389925 contigs are shown in the table below, along with their locations relative to SEQ ID NOs 19, 20, 22, and 24; note that the 5' and 3' untranslated regions may extend further along the contig sequence beyond those portions that correspond to SEQ ID NOs 19, 20, 22, and 24, as indicated by the parentheses around the AL109843 and AL389925 endpoints in the table. Due to the preliminary nature of the sequence data and assembly of the contig sequence, the exons within the genomic contigs may contain sequence variations due to inaccurate sequence data or allelic polymorphism.

Corresponding positions of HPR2 gene exons in human genomic contigs AL109843 and AL389925 and in HPR2 coding sequences:

	Position in AL109843	Position in SEQ ID NO:19 / 20 / 22 / 24
Exon 1	(34088)-34164	1-77 / (5' UTR, not in SEQ ID NOs 20, 22, and 24)
Exon 2	35715-35813	78-176 / 1-70 / 1-70 / 1-70
Exon 3	36965-37261	177-473 / 71-367 / 71-367 / 71-367
Exon 4	50459-50582	474-597 / 368-491 / 368-491 / 368-491
Exon 5	68360-68520	598-758 / 492-652 / 492-652 / 492-652
Exon 6	74533-74678	759-904 / 653-798 / 653-798 / 653-798
Exon 7	87197-87353	905-1061 / 799-955 / 799-955 / 799-955
Exon 8	104336-104425	1062-1151 / 956-1045 / (not present) / 956-1045
Exon 9	107802-107904	1152-1254 / 1046-1148 / (not present) / (not present)
	Position in AL389925	Position in SEQ ID NO:19 / 20 / 22 / 24
Exon 10	8847-8937	1255-1345 / 1149-1239 / 'G'-957-1047 / 1046-1071
Exon 11	11488-(12972)	1346-2830 / 1240-1890 / 1048-1698 / (not present)

In the HPR3-ex9 splice variant, note that the absence of the exon 9 sequence (103 nucleotides) changes the reading frame towards the 3' end of the coding sequence for the HPR2-ex9 form (SEQ ID NO:24)

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relative to that of the HPR2 coding sequence of SEQ ID NO:20, leading to a different amino acid sequence in the HPR2-ex9 C-terminal portion and a stop codon after amino acid 356 (compared to 629 amino acids in HPR2). For the HPR2-ex8-ex9 form, the splice is made at a slightly different exon 10 splice acceptor site than for the HPR2 form, so that an extra 'G' residue is included at the start of exon 10 in the HPR2-ex8-ex9 form, restoring the reading frame to be the same as in the 3' end of the HPR2 sequence. The C-terminal 248 amino acids of HPR2-ex8-ex9 form are therefore the same as the C-terminal 248 amino acids of HPR2 form, and although the coding sequence of the HPR2-ex8-ex9 form is missing both exons 8 and 9 (except for the last 'G' residue of exon 9), the resulting HPR2-ex8-ex9 form polypeptide is longer (565 amino acids) than the HPR2-ex9 form polypeptide (356 amino acids).

Several splice variations of the HPR2 sequences have been identified in human genomic sequences and are included within the scope of the invention. For example, amino acids 118 through 215 of SEO ID NO:16 match the amino acid sequence of HPR2 presented in SEQ ID NO:21, while amino acids 1 through 117 of SEQ ID NO:16 may correspond to an alternatively spliced exon added upstream of exon 3 (i.e. at the exon/intron boundary identified between nucleotides 176 and 177 of SEO ID NO:19). Amino acids 216 through 245 of SEQ ID NO:16 may correspond to an additional alternatively spliced exon added between exon 3 and exon 4 (i.e. at the exon/intron boundary identified between nucleotides 473 and 474 of SEQ ID NO:19). Amino acids 340 through 344 of SEQ ID NO:16 may correspond to an alternatively spliced exon added downstream of exon 5 (i.e. at the exon/intron boundary identified between nucleotides 758 and 759 of SEQ ID NO:19). In a further potential splice variant, an alternative exon or exons encoding the amino acid sequence of SEQ ID NO:17 could be substituted for exon 6, resulting in the replacement of amino acids 217 through 267 of SEQ ID NO:21 with the SEQ ID NO:17 amino acids. In this potential variant, 51 amino acids N-terminal to the WSXWS motif, including the proline-rich region (as shown in Table 1) between the two cytokine receptor subdomains, would be replaced with 39 amino acids, resulting in deletion of a portion of the more C-terminal cytokine receptor subdomain which includes a highly conserved Trp residue. In an additional potential splice variant, an alternative exon could be added downstream of exon 4 (i.e. at the exon/intron boundary identified between nucleotides 597 and 598 of SEQ ID NO:19) so that an amino acid sequence starting in the amino acids of SEQ ID NO:18 could be substituted for amino acids following and including the serine at position 164 of SEQ ID NO:21. Multiple splice variations as described above can be included in a single splice variant, for example, replacing exon 6 with an alternative exon or exons encoding the amino acid sequence of SEQ ID NO:17, and also deleting exons 8 and/or 9 as described above.

Additional variations of HPR2 polypeptides are provided as naturally occurring genomic variants of the HPR2 sequences disclosed herein; such variations may be incorporated into an HPR2 polypeptide or nucleic acid individually or in any combination, or in combination with alternative splice variation as described above. As one example, a change from the Leu residue position 310 of SEQ ID NO:21 to a Pro residue could be caused by a single change from 'T' to 'C' at position 1035 of SEO ID NO:19. This variation and another are listed in the table below:

Amino Acid Change	Position in SEQ ID NO:21	Nucleotide Change	Position in SEQ ID NO:19
Leu->Pro	310	T->C	1035
(not applicable)	(not applicable)	A->G	2172 (3' UTR)

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A nucleic acid encoding a polypeptide with a high degree of amino acid similarity (approximately 69% amino acid identity) to human HPR2 was isolated from Mus musculus. The Mus HPR2 amino acid sequence is presented as SEQ ID NO:27, and due to its high level of similarity with human HPR2, is considered to be the murine homologue of human HPR2. PCR amplification of cDNA sequences corresponding to mRNAs encoding murine HPR2 identified a cDNA molecule encoding SEQ ID NO:27; the nucleotide sequence of this murine HPR2 cDNA is presented as SEQ ID NO:29. Nucleotides 1 through 1932 of SEQ ID NO:29 encode SEQ ID NO:27, with nucleotides 1933-1935 corresponding to a stop codon. The murine HPR2 amino acid sequence of SEQ ID NO:27 appears to have a 20-amino acid insertion at amino acids 297 through 316 of SEQ ID NO:27 relative to human HPR2 of SEQ ID NO:21, based on an alignment of the human and murine polypeptide sequences; this insertion is identical to amino acids 317 through 336. Given the number of alternatively spliced forms identified for human HPR2, it is possible that this insertion in murine HPR2 relative to the human HPR2 of SEQ ID NO:21 is the result of alternative splicing. One embodiment of the invention is a form of murine HPR2 in which one of these repeated WQPWS-containing motifs has been deleted; that is, polypeptides in which the amino acid sequence ending with amino acid 296 of SEQ ID NO:27 is contiguous with the amino acid sequence beginning with amino acid 317 of SEQ ID NO:27, or polypeptides in which the amino acid sequence ending with amino acid 316 of SEQ ID NO:27 is contiguous with the amino acid sequence beginning with amino acid 337 of SEQ ID NO:27.

## C. Comparison of HPR1 and HPR2 to Other Hematopoietin Receptor Polypeptides.

The amino acid sequences of human HPR1 (SEQ ID NO:4), murine HPR1 (SEQ ID NO:12), and human HPR2 (SEQ ID NO:21) were compared with the amino acid sequences of these other hematopoietin receptor family members - LIF-R, the interleukin 12 beta 2 receptor chain (IL-12RB2), gp130, and GCSFR (SEQ ID NO:6 - SEQ ID NO:9, respectively) - using the GCG "pretty" multiple sequence alignment program, with amino acid similarity scoring matrix = blosum62, gap creation penalty = 8, and gap extension penalty = 2. Alignments of these sequences are shown in Table 1, and include consensus residues which are identical among at least three of the amino acid sequences in the alignment. The capitalized residues in the alignment are those which match the consensus residues. The numbering of amino acid residues in Table 1 corresponds to the position of those residues in the HPR1 amino acid sequence (SEQ ID NO:4). Note that only a portion of the HPR2 amino acid sequence is shown in Table 1, as HPR2 does not contain fibronectin type III repeats in its extracellular domain. HPR1 and HPR2 sequences corresponding to the intracellular Box 1 and Box 2 motifs are shown in Table 2. Sequences of eleven amino acids similar to the Box 1 or 2 motif of other hematopoietin receptors were identified for HPR1 and HPR2, and placed into a column with these motif sequences (with no gaps introduced). Similarly, HPR2 sequences corresponding to the

5 intracellular Box 3 motif are shown in Table 3. Sequences of fourteen amino acids similar to the Box 3 motif of other hematopoietin receptors were identified for HPR2, and placed into a column with these motif sequences (with no gaps introduced). The numbering of each sequence on Tables 2 and 3 corresponds to their position in the complete amino acid sequence for that HPR polypeptide. The consensus residues are those that are present in three or more (for Table 2) or two or more (for Table 3) sequences at that position in the motif.

Amino acid substitutions and other alterations (deletions, insertions, etc.) to HPR1 and HPR2 amino acid sequences (for example, SEQ ID NOs 4, 12, and 21) are predicted to be more likely to alter or disrupt HPR1 or HPR2 polypeptide activities if they result in changes to the capitalized residues of the amino acid sequences as shown in Tables 1, 2, and 3, and particularly if those changes do not substitute an amino acid of similar structure (such as substitution of any one of the aliphatic residues -Ala, Gly, Leu, Ile, or Val - for another aliphatic residue), or a residue present in other hematopoietin receptor polypeptides at that conserved position. Conversely, if a change is made to an HPR1 or HPR2 amino acid sequence resulting in substitution of the residue at that position in the alignment from one of the other Table 1, 2, or 3 hematopoietin receptor polypeptide sequences, it is less likely that such an alteration will affect the function of the altered HPR1 or HPR2 polypeptide. For example, the consensus residue at position 42 in Table 1 is serine, and one of the hematopoietin receptors (LIF-R) has an asparagine at that position. Substitution of asparagine or the chemically similar glutamine for serine at that position is considered to be less likely to alter the function of the polypeptide than substitution of tryptophan or tyrosine etc. Embodiments of the invention include HPR1 and HPR2 polypeptides and fragments of HPR1 and HPR2 polypeptides, comprising altered amino acid sequences. Altered HPR1 or HPR2 polypeptide sequences share at least 30%, or more preferably at least 40%, or more preferably at least 50%, or more preferably at least 55%, or more preferably at least 60%, or more preferably at least 65%, or more preferably at least 70%, or more preferably at least 75%, or more preferably at least 80%, or more preferably at least 85%, or more preferably at least 90%, or more preferably at least 95%, or more preferably at least 97.5%, or more preferably at least 99%, or most preferably at least 99.5% amino acid identity with one or more of the hematopoietin receptor amino acid sequences shown in Tables 1, 2, and 3.

Table 1: Alignment of HPR1 and HPR2 extracellular domains with those of other hematopoietin receptors

35 C: conserved cysteine

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proline-rich 'linker' between cytokine receptor subdomains

: WSXWS motif

i fibronectin type III repeatsi transmembrane domain

	SEQ ID		
	NO:	35	3
Hs HPR1	4	PaKPeNiSCV yYyr.KNLTC TWsPGkETsyTqYTv KrtyafGekh	ı
Mus HPR1	12	PtKPeNiSCV fYfd.rNLTC TWrPekETndTSYiv tltySyGK	
gp130	8	PeKPkNLSCi vneg.KkmrC eWdgGrETHLeTnfTL KsewathKfa	
GCSFR	9	PaiPhNLSCl mnlttssLiC qWePGpETHLpTSfTL KsfkSrGnCq	1
Hs HPR2	21	PdiPdevtCV iYeysgNmTC TWnaGklTyidTkYvv hvKsl	L

IL-12RB2	7	PeqPqNLS <b>C</b> i	qkgeqgtva <b>C</b>	TWerGrdTHL	$\dots$ yTeYTL	qlSgpKnl
LIF-R	6	Pdt PaaInCe	th.dlKeii <b>C</b>	sWnPGrvTaL	vgpraTSYTL	vesfS.GKvv
	<u> </u>	D_KD_MT.CCV	-VKNT.TC	TW-PG-ETHL	TSYTI	KS-GK
consensus		F-KF-NDSCV_	1 - 101110	177 1 0 111111	10111	10 B G10
		79				126
Hs HPR1	4	dn <b>C</b> ttnssts	enras <b>C</b> sffL	PRitipdN	YtieVeAENg	dGvikShmty
Mus HPR1	12	snys	dnateasysf	PRscamppdi	csVeVQAqNg	dGkvkSdity
gp130	8	d <b>C</b> kakr	dtptsCtvdv	stvyfvN	ieVWVeAENA	LGkvtSdhin
	9	+codeilder	nkdagah <b>C</b> ai	PRkhlllyqN	maiwwww.ema	T.Ct SmSncT.c
GCSFR		tagasiracv	pragasiici		MIGINACALINA	T.Cman.Clast.co
Hs HPR2	21_	ereeeddarr	ssyini	stdsląggkk	YIVWVQAANA	LGMeeskqLq
IL-12RB2	7	twqkq <b>C</b> kdiy	<b>C</b> dYldfginL	tpespesN	<u>ttakVtAvNs</u>	LGSSSSIpst
LIF-R	6	rlkraeaptn	esYqllfqmL	Pnqei	YnftlnAhNp	LGrSqStiL.
consensus			YL	PRN	Y-VWVOAENA	LG-S-SL-
		127				172
			DD1-T CDIVID	7	i Oi eWileDell	
Hs HPR1	4	wrleniakte	PPKIERVKPV	lgikrm	TOTEMIKEET	apvssdLKyt
Mus HPR1	12	whLisIaKtE	PPiIlsVnPI	cnrm	fQiqW.kPre	ktrgfpLvCm
gp130_	8	fdpvykVKPn	PPhnlsVins	eelssi	lkLtWtnPsi	ksv.IiLKyn
GCSFR	9			peaappqagc		pglhInqKCE
Hs HPR2	21	ihLdDTViPa	aavTsRaetT	natvpkti	ivWds	
IL-12RB2	7	f+flntv~n1	DDWdiDivf~	kasvsrct	Tariated	eglvLlnr
LIF-R	6			nsta		nfakInflCE
consensus		L-DIVKPE	PP-I-RVKPI		-QL-WP	I-LKCE
	1					
		173				221
Hs HPR1	4		sWmeVnFakN	rkdknqtynL	tGLabetevi	
Mus HPR1	12			ckqvcnL		
gp130	8	iqYRTkda.s	twsqip.ped	tastrssftv	darkbt.l.e.k.	FrikCmkEag
GCSFR	9	LRhkpqrgea	sWalVgp	lplealqyeL	cGLlPaTaYt	lQIRCirwpl
Hs HPR2	21	mRYkattngt	.WnvkeFd	tnftyvqqse	fyLePnikYV	FQvRCq.Etg
IL-12RB2	7			vtkakgrhdL		
T.TF_P	6	ieikkeNSva	earnVti ka			FrIRCstEtf
LIF-R	6			venssylvaL	dkLnPyTlYt	
LIF-R consensus	6				dkLnPyTlYt	
	6			venssylvaL	dkLnPyTlYt	
	6	LRYRT-NS		venssylvaL	dkLnPyTlYt	FQIRCE
	6	LRYRT-NS	-WV-FN	venssylvaL L	dkLnPyT1Yt -GL-PFTEYV	FQIRCE 261
consensus		LRYRT-NS	-WV-FN	venssylvaL L	dkLnPyT1Yt -GL-PFTEYV	FQIRCE 261
Consensus  Hs HPR1	4	LRYRT-NS 222 K.fWSDWSqE	-WV-FN kmgmTeEEaP	venssylvaL L cgLe	dkLnPyTlYt -GL-PFTEYV lWRvLkP	FQIRCE 261 aeadGrRpVr
Hs HPR1 Mus HPR1	4 12	LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE	-WV-FN kmgmTeEEaP etrvTmEEvP	venssylvaLL cgLe hvLD	dkLnPyTlYt -GL-PFTEYV  lwRvLkP lwRiLeP	261 aeadGrRpVr admnGdRkVr
Hs HPR1 Mus HPR1 gp130	4 12 8	LRYRT-NS 222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE	-WV-FN kmgmTeEEaP etrvTmEEvP asgiTyEdrP	cgLe hvLD skaps	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP	261 aeadGrRpVr admnGdRkVr shtqGyRtVq
Hs HPR1 Mus HPR1 gp130 GCSFR	4 12 8 9	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP	cgLe hvLD skaps tvrLD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP	261 aeadGrRpVr admnGdRkVr shtqGyRtVq
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2	4 12 8 9 21	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSs1	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP	cgLe hvLD skaps tvrLD qvtskafqhD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq
Hs HPR1 Mus HPR1 gp130 GCSFR	4 12 8 9	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSs1	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP	cgLe hvLD skaps tvrLD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2	4 12 8 9 21	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSs1 KGSWSDWSes	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP	cgLe hvLD skaps tvrLD qvtskafqhDtgmlD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD.	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R	4 12 8 9 21 7	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSs1 KGSWSDWSes .wkWSkWSnk	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP	cgLe hvLD skaps tvrLD qvtskafqhDtgmlD skgpD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknli
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2	4 12 8 9 21 7	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSs1 KGSWSDWSes .wkWSkWSnk	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~ysrqqisssdgknliG-R-V-
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R	4 12 8 9 21 7	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSs1 KGsWSDWSes .wkWSkWSnk KGYWSDWS-E	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~ysrqqisssdgknliG-R-V-
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSs1 KGsWSDWSes .wkWSkWSnk KGYWSDWS-E	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP	cgLe hvLD skaps tvrLD qvtskafqhDtgmlD skgpD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~ysrqqisssdgknliG-R-V-
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP T-EE-P	cgLe hvLD skaps tvrLD qvtskafqhDtgmlD skgpDLD	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~ysrqqisssdgknliG-R-V-
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP T-EE-P	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhDtgmlD skgpDLD wyypesnTn. qyfaensTn.	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~ysrqqisssdgknliG-R-V-
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus HS HPR1 Mus HPR1	4 12 8 9 21 7 6	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP T-EE-P	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhDtgmlD skgpDLD wyypesnTn. qyfaensTn.	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus  HS HPR1 Mus HPR1 gp130	4 12 8 9 21 7 6	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LvWKtLPpfE	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP T-EE-P vleKtLGYni vleKtfGYhi AnGKILdYeV	venssylvaLL  cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl.	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~ysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus  HS HPR1 Mus HPR1 gp130 GCSFR	4 12 8 9 21 7 6 4 12 8 9	LRYRT-NS  222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LvWKtLPpfE LfWKpvPleE	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus  HS HPR1 Mus HPR1 gp130 GCSFR IL-12RB2	4 12 8 9 21 7 6 4 12 8 9	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpVPleE LfWKnLsvsE	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP T-EE-P vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus  HS HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R	4 12 8 9 21 7 6 4 12 8 9	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpVPleE LfWKnLsvsE iyWKpLPinE	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV AnGKILsYnV	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqs	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdknd
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus  HS HPR1 Mus HPR1 gp130 GCSFR IL-12RB2	4 12 8 9 21 7 6 4 12 8 9	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGsWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpvPleE LfWKpvPleE LfWKpvPleE LfWKpLFine L-WK-LPE	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV AnGKILsYnV	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdknd
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus  HS HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R	4 12 8 9 21 7 6 4 12 8 9	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSeE pGhWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpvPleE LfWKpvPleE LfWKpvPleE LfWKpLFinE L-WK-LP-E	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV AnGKILsYnV	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqs	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL
HS HPR1 Mus HPR1 gp130 GCSFR HS HPR2 IL-12RB2 LIF-R consensus  HS HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R	4 12 8 9 21 7 6 4 12 8 9	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGsWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpvPleE LfWKpvPleE LfWKpvPleE LfWKpLFine L-WK-LPE	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV AnGKILsYnV	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqs	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdknd
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6 4 12 8 9 7 6	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGsWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpVPleE LfWKpVPleE LfWKpVPleE LfWKpVPleE LfWKpVPleE LFWKPLP-E	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasP T-EE-P vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV AnGKILSYnV A-GKILGY-V	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqsT	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~~~ysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6 4 12 8 9 7 6	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGsWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpVPleE	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV ArGKILhYqV ArGKILhYqV ArGKILSYnV A-GKILGY-V	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqsT	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~~ysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6 4 12 8 9 7 6	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGsWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpLPinE LfWKpLPinE L-WK-LPE  311 fwVSmisyNS hsVSVtsfNS	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILhYqV AGKILSYNV A-GKILGY-V	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqsT IPaiqEksfq IPdvhEktfq	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T cievmqAcva yiksmqAyia	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~~ysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.1LvVnWQ
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6 4 12 8 9 7 6	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpVPleE LF	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV ArGKILhYqV ArGKILhYqV ArGKILSYnV A-GKILGY-V  IGKSpvatLr lGKSqeTiLr vGKSdaavLt	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqsT IPaiqEksfq IPdvhEktfq IPacdfqath	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T cievmqAcva yiksmqAyia pvmdlkAfpk	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~~~ysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL \$359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6 4 12 8 9 7 6	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGsWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LVWKtLPpfE LfWKpvPleE LfWKpvPleS jyWKpLPinE L-WK-LPE 311 fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILhYqV ArGKILSYNV A-GKILGY-V lGKSpvatLr lGKSqeTiLr vGKSdaavLt aGtSrPTpv.	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqsT IPaiqEksfq IPacdfqath .vfsEsrgp	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT. ilplcNtTel mTqnitghts LSEipd.pqh LTE-N-T- cievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVq ~~~~~~~ysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdknd
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6 4 12 8 9 7 6	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LlWKkLPpfE LfWKpvPleE LfWKpvPleE iyWKpLPinE L-WK-LPE  311 fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS waVaVsAaNS	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtLGYni vleKtLGYni vleKtLGYni vleKtLGYni AngKILdyeV dsGrigGyvV ArgKiLhyqV ArgKiLhyqV ArgKiLhyqv lGKSpvatLr lGKSpvatLr lGKSqeTiLr vGKSdaavLt agtSrPTpv. kGsSlPTrin	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgaga tlqeltggka scssdeeTqsT IPaiqEksfq IPdvhEktfq IPacdfqath .vfsEsrgp ImnlcEagll	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T- cievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar aprqvsAnse	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wtviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt dphsLwVgWe gmdnilVtWQ
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6 4 12 8 9 7 6	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LlWKkargap LlWKkLPpfE LfWKpLPinE LfWKpLPinE LfWK-LPE  311 fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS waVaVsAaNS yiiSVvAkNS	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILSYnv A-GKILGY-V  lGKSpvatLr lGKSqeTiLr vGKSdaavLt aGtSrPTpv. kGsSlPTrin vGsSpPskia	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqsT IPaiqEksfq IPdvhEktfq IPacdfqath .vfsEsrgp ImnlcEagll smeipnddl.	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T- cievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar aprqvsAnsekieqvvg	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt dphsLwVgWe gmdnilVtWQ mgkgilltWh
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6 4 12 8 9 7 6	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LlWKkargap LlWKkLPpfE LfWKpLPinE LfWKpLPinE LfWK-LPE  311 fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS waVaVsAaNS yiiSVvAkNS	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILSYnv A-GKILGY-V  lGKSpvatLr lGKSqeTiLr vGKSdaavLt aGtSrPTpv. kGsSlPTrin vGsSpPskia	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqsT IPaiqEksfq IPdvhEktfq IPacdfqath .vfsEsrgp ImnlcEagll smeipnddl.	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T- cievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar aprqvsAnsekieqvvg	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt dphsLwVgWe gmdnilVtWQ mgkgilltWh
Hs HPR1 Mus HPR1 gp130 GCSFR Hs HPR2 IL-12RB2 LIF-R consensus  Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2 LIF-R consensus	4 12 8 9 21 7 6 4 12 8 9 7 6	222 K.fWSDWSqE r.YWSkWSkE KGYWSDWSps KrYWqpWSsl KGSWSDWSes .wkWSkWSnk KGYWSDWS-E  262 LlWKkargap LlWKkargap LlWKkargap LlWKkLPpfE LfWKpLPinE LfWKpLPinE LfWK-LPE  311 fwVSmisyNS hsVSVtsfNS ylatltvrNl qeValvAyNS waVaVsAaNS yiiSVvAkNS	kmgmTeEEaP etrvTmEEvP asgiTyEdrP lelrTtEraP ffhkTpEtvP lraqTpEEeP kqhlTtEasPT-EE-P  vleKtLGYni vleKtfGYhi AnGKILdYeV dsGrIqGYvV ArGKILhYqV ArGKILhYqV AnGKILSYnv LGKSQeTiLr vGKSQavLt aGtSrPTpv. kGsSlPTrin vGsSpPskia -GKS-PT-L-	cgLe hvLD skaps tvrLD qvtskafqhD .tgmlD skgpDLD  wyypesnTn. qyfaensTn. qyfaensTn. tltrwkshl. swrpsgqaga tlqeltggka scssdeeTqsT IPaiqEksfq IPdvhEktfq IPacdfqath .vfsEsrgp ImnlcEagll smeipnddl. IPE	dkLnPyTlYt -GL-PFTEYV  lWRvLkP lWRiLeP fWykiDP TWwrqRqLDP TWns~~~~~ vWymkRhiD. TWrew TWR-LDP  LTEtmNtTnq LTEinNiTtq qnytvNaT. ilplcNtTel mTqnitghts LsEipd.pqh LTE-N-T- cievmqAcva yiksmqAyia pvmdlkAfpk altrlhAmar aprqvsAnsekieqvvgA	261 aeadGrRpVr admnGdRkVr shtqGyRtVqRtVqysrqqisssdgknliG-R-V- 310 qlelhLgges qyellLmsqa kltvnLtndr sctfhLpsea wttviprtgn kaeirLdkndL 359 ed.qLvVkWQ ep.lLvVnWQ dn.mLwVeWt dphsLwVgWe gmdnilVtWQ mgkgilltWh

		360				404
Hs HPR1	4	ssaldVn t	wmIEWfpdv	d.SePttlsW	e.svSqaTnw	TIqqDkLKPF
Mus HPR1	12	ssipaVd t				TIeqDkLKPF
gp130	8	tPresVk k				TylrgNLaes
GCSFR	9	pPnpwpq g				fllkeNirPF
IL-12RB2	7	pPrkdpsaVq e	YVvEWrelh	pggdtqvpln	wlrsrpynvs	aliseNiKsy
LIF-R	6	ydpnmtc d	lYVIkWc.ns	srSePclmdW	rkvpSnsTet	vIesDefrPg
consensus		-PV	YVIEW	S-PW	ST	TIDNLKPF
		405				454
Hs HPR1	4	wCYNIsVYPm 1	hDkvGePyS	IqAYaKEgvP	SeGPEtkVEn	IGvktvtItW
Mus HPR1	12			IqAYaKEgtP	lkGPEtrVEn	IGlrtAtItW_
gp130	8			IkAYlKqapP	SkGPtvrtkk	vGKneAvleW
GCSFR	9			vyAYsqEmAP	ShaPElhlkh	IGKtwAqleW
IL-12RB2	7			IlgnsKhkAP	lsGPhinait	eeKgsilIsW
LIF-R	6	irYNfflYgc r	nggyqllrS	migYieElAP	ivaPnftVEd	tsadsilvkW
consensus		-CYNI-VYP	-DG-P-S	I-AY-KE-AP	S-GPEVE-	IGKA-I-W
		455				489
Hs HPR1	4	keIPksErkG i			<u></u>	.VNSSi
Mus HPR1	12			aeGGkelSKt		.VNSha
gp130	8					.VdSSh
GCSFR	9	vpePpelgks p				.lNaSs
IL-12RB2	7	nsIPVqEqmG c				.ceipyrvSq
LIF-R	6	edIPVeElrG F	rlray.lFYf	akGerdtSKm	rylesarsdi	kVknitdiSq
consensus		IPV-EG F				-VNSS-
consensus		IPV-EG F				-VNSS-
		IPV-EG F	NYTIFY-	GGSK-		-VNSS- 539
Hs HPR1	4	490 lqygLeSLkr k	TNYTIFY-	GGSK- sTsAGGTNGt	sinFkTLsfS	-VNSS- 539 VfEIilltsL
	12	490 lqygLeSLkr k lqcdLeSLtr r	TNYTIFY- TSYiVqvMA TSYtVwvMA	GGSK- sTsAGGTNGt sTrAGGTNGV	sinFkTLsfS rinFkTLsiS	-VNSS- 539 VfEIilltsL VfEvvlltsL
Hs HPR1	12	490 lqygLeSLkr k lqcdLeSLtr r	TNYTIFY- TSYiVqvMA TSYtVwvMA	sTsAGGTNGt sTrAGGTNGV yTdeGGkdGp	sinFkTLsfS rinFkTLsiS eftFtTpkfa	-VNSS- 539 VfEIilItsL VfEvvlltsL qgEIeaIvvp
Hs HPR1 Mus HPR1 gp130 GCSFR	12 8 9	490 lqygLeSLkr k lqcdLeSLtr r teytLsSLts d rgfvLhgLep a	T-NYTIFY- TSYIVQVMA TSYtVwvMA TIYmVrmaA aslyhihlMA	sTsAGGTNGt sTrAGGTNGv yTdeGGkdGp asqAGaTNst	sinFkTLsfS rinFkTLsiS eftFtTpkfa vltlmTLtpe	-VNSS- 539 VfEIilItsL VfEvvlltsL qgEIeaIvvp gsElhiIlgL
Hs HPR1 Mus HPR1 gp130	12 8 9 7	490 lqygLeSLkr k lqcdLeSLtr r teytLsSLts d rgfvLhgLep a nshpinSLqp r	T-NYTIFY- TSYiVqvMA TSYtVwvMA TIYmVrmaA aslyhihlMA rvtyvlwmtA	sTsAGGTNGt sTrAGGTNGv yTdeGGkdGp asqAGaTNst lTaAGesshg	sinFkTLsfS rinFkTLsiS eftFtTpkfa vltlmTLtpe nerefcLqgk	539 VfEIilItsL VfEvvlltsL qgEIeaIvvp gsElhiIlgL anwmafvaps
Hs HPR1 Mus HPR1 gp130 GCSFR	12 8 9	490 lqygLeSLkr k lqcdLeSLtr r teytLsSLts d rgfvLhgLep a nshpinSLqp r ktlriadLqg k	T-NYTIFY- TSYIVQVMA TSYtVwvMA TIYmVrmaA aslyhihlMA rvtYvlwmtA TSYhlvlrA	sTsAGGTNGt sTrAGGTNGv yTdeGGkdGp asqAGaTNst lTaAGesshg yTdgGvgpek	sinFkTLsfS rinFkTLsiS eftFtTpkfa vltlmTLtpe nerefcLqgk smyvvTkenS	539 VfEIilItsL VfEvvlltsL qgEIeaIvvp gsElhiIlgL anwmafvaps VglIiaIlip
Hs HPR1 Mus HPR1 gp130 GCSFR IL-12RB2	12 8 9 7	490 lqygLeSLkr k lqcdLeSLtr r teytLsSLts d rgfvLhgLep a nshpinSLqp r ktlriadLqg k	T-NYTIFY- TSYIVQVMA TSYtVwvMA TIYmVrmaA aslyhihlMA rvtYvlwmtA TSYhlvlrA	sTsAGGTNGt sTrAGGTNGv yTdeGGkdGp asqAGaTNst lTaAGesshg	sinFkTLsfS rinFkTLsiS eftFtTpkfa vltlmTLtpe nerefcLqgk smyvvTkenS	539 VfEIilItsL VfEvvlltsL qgEIeaIvvp gsElhiIlgL anwmafvaps VglIiaIlip

Table 2: Box 1 and Box 2 motifs in the intracellular domains of HPR1, HPR2, and other hematopoietin receptors

	SEQ ID NO	Box 1 Motif	Box 2 Motif
Hs HPR1	4	563-thlcWPtV <b>P</b> NP-573	631-eifTdEArtgq-641
Mus HPR1	12	517-tplccPDVPNP-527	582-VvlTEEAgKgq-592
HPR2	21	393-pkwlyeDi <b>P</b> Nm-403	430-VdpmiteiKei-440
LIF-R	6	866-KetfyPDi <b>P</b> NP-876	910-VleTrsAfpKi-920
gp130	8	648-KkhiWPnV <b>P</b> dP-658	693-VveiEandKKp-703
GCSFR	9	655-KnplWPsV <b>P</b> dP-665	696-ltvlEEdeKKp-706
consensus		KWPDV <b>P</b> NP	VTEEA-KK-

10 Table 3: Box 3 motifs in the intracellular domains of HPR2 and other hematopoietin receptors

	SEQ ID NO	Box 3 Motif
HPR2 (first occurrence)	21	478-PdLntGYKPQisnf-491
HPR2 (second occurrence)	21	605-lpsint <b>Y</b> fP <b>Q</b> niLe-618
LIF-R	6	995-PVggaGYKPQmhLp-1008
gp130	8	693-tVvhsGYrhQvpsv-774
GCSFR	9	696-PtLvqt <b>Y</b> vl <b>Q</b> gdpr-734
consensus residues		PVLG <b>Y</b> KP <b>Q</b> L-

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# **EXAMPLE 2: Monoclonal Antibodies That Bind Polypeptides of the Invention**

This example illustrates a method for preparing monoclonal antibodies that bind HPR1 or HPR2 polypeptides. Suitable immunogens that may be employed in generating such antibodies include, but are not limited to, purified HPR1 or HPR2 polypeptide or an immunogenic fragment thereof. Purified HPR1 or HPR2 polypeptide can be used to generate monoclonal antibodies immunoreactive therewith, using conventional techniques such as those described in U.S. Patent 4,411,993. Briefly, mice are immunized with HPR1 or HPR2 polypeptide immunogen emulsified in complete Freund's adjuvant, and injected in amounts ranging from about 10 to about 100 micrograms subcutaneously or intraperitoneally. Ten to twelve days later, the immunized animals are boosted with additional HPR1 or HPR2 polypeptide emulsified in incomplete Freund's adjuvant. Mice are periodically boosted thereafter on a weekly to bi-weekly immunization schedule. Serum samples are periodically taken by retro-orbital bleeding or tail-tip excision to test for anti-HPR1 or anti-HPR2 antibodies by dot blot assay, ELISA (Enzyme-Linked Immunosorbent Assay), or inhibition of binding of HPR1 or HPR2 polypeptide to an HPR1 and/or HPR2 binding partner.

Following detection of an appropriate antibody titer, positive animals are provided one last intravenous injection of HPR1 or HPR2 polypeptide in saline. Three to four days later, the animals are sacrificed, spleen cells harvested, and spleen cells are fused to a murine myeloma cell line, e.g., NS1 or preferably P3x63Ag8.653 (ATCC CRL 1580). Fusions generate hybridoma cells, which are plated in multiple microtiter plates in a HAT (hypoxanthine, aminopterin and thymidine) selective medium to inhibit proliferation of non-fused cells, myeloma hybrids, and spleen cell hybrids.

The hybridoma cells are screened by ELISA for reactivity against purified HPR1 or HPR2 polypeptide by adaptations of the techniques disclosed in Engvall et al., (*Immunochem. 8*:871, 1971) and in U.S. Patent 4,703,004. A preferred screening technique is the antibody capture technique described in Beckmann et al., (*J. Immunol. 144*:4212, 1990). Positive hybridoma cells can be injected intraperitoneally into syngeneic BALB/c mice to produce ascites containing high concentrations of anti-HPR1 or anti-HPR2 monoclonal antibodies. Alternatively, hybridoma cells can be grown *in vitro* in flasks or roller bottles by various techniques. Monoclonal antibodies produced in mouse ascites can be purified by ammonium sulfate precipitation, followed by gel exclusion chromatography. Alternatively, affinity chromatography based upon binding of antibody to Polypeptide A or Polypeptide G can also be used, as can affinity chromatography based upon binding to HPR1 or HPR2 polypeptide.

### **EXAMPLE 3**

### Antisense Inhibition of HPR1 and/or HPR2 Expression

In accordance with the present invention, a series of oligonucleotides are designed to target different regions of HPR1 and/or HPR2 human or murine mRNA molecules, using the nucleotide sequences of SEQ ID NOs 3, 5, 19, 20, 22, 24, 28, and 29 as the bases for the design of the oligonucleotides. The oligonucleotides are selected to be approximately 10, 12, 15, 18, or more

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preferably 20 nucleotide residues in length, and to have a predicted hybridization temperature that is at least 37 degrees C. Preferably, the oligonucleotides are selected so that some will hybridize toward the 5' region of the mRNA molecule, others will hybridize to the coding region, and still others will hybridize to the 3' region of the mRNA molecule.

The oligonucleotides may be oligodeoxynucleotides, with phosphorothioate backbones (internucleoside linkages) throughout, or may have a variety of different types of internucleoside linkages. Generally, methods for the preparation, purification, and use of a variety of chemically modified oligonucleotides are described in U.S. Patent No. 5,948,680. Modified oligonucleosides may also be used in oligonucleotide synthesis, as well as mixed backbone compounds having, for instance, alternating MMI and P=O or P=S linkages, which are prepared as described in U.S. Pat. Nos. 5,378,825, 5,386,023, 5,489,677, 5,602,240 and 5,610,289. Formacetal- and thioformacetal-linked oligonucleosides may also be used and are prepared as described in U.S. Pat. Nos. 5,264,562 and 5,264,564; and ethylene oxide linked oligonucleosides may also be used and are prepared as described in U.S. Pat. No. 5,223,618. Peptide nucleic acids (PNAs) may be used as in the same manner as the oligonucleotides described above, and are prepared in accordance with any of the various procedures referred to in Peptide Nucleic Acids (PNA): Synthesis, Properties and Potential Applications, Bioorganic & Medicinal Chemistry, 1996, 4, 5-23; and U.S. Pat. Nos. 5,539,082, 5,700,922, and 5,719,262. Chimeric oligonucleotides, oligonucleosides, or mixed oligonucleotides/oligonucleosides of the invention are synthesized according to U.S. Pat. No. 5,623,065.

The effect of antisense compounds on target nucleic acid expression can be tested in any of a variety of cell types provided that the target nucleic acid is present at measurable levels. This can be routinely determined using, for example, PCR or Northern blot analysis. Preferably, the effect of several different oligonucleotides should be tested simultaneously, where the oligonucleotides hybridize to different portions of the target nucleic acid molecules, in order to identify the oligonucleotides producing the greatest degree of inhibition of expression of the target nucleic acid. Antisense modulation of HPR1 and/or HPR2 nucleic acid expression can be assayed in a variety of ways known in the art. For example, HPR1 and HPR2 mRNA levels can be quantitated by, e.g., Northern blot analysis, competitive polymerase chain reaction (PCR), or real-time PCR (RT-PCR). Real-time quantitative PCR is presently preferred. RNA analysis can be performed on total cellular RNA or poly(A)+ mRNA. Methods of RNA isolation and Northern blot analysis are taught in, for example, Ausubel, F. M. et al., Current Protocols in Molecular Biology, Volume 1, pp. 4.1.1-4.2.9 and 4.5.1-4.5.3, John Wiley & Sons, Inc., 1996. Real-time quantitative (PCR) can be conveniently accomplished using the commercially available ABI PRISM 7700 Sequence Detection System, available from PE-Applied Biosystems, Foster City, Calif. and used according to manufacturer's instructions. Other methods of quantitative PCR analysis are also known in the art. HPR1 and HPR2 protein levels can be quantitated in a variety of ways well known in the art, such as immunoprecipitation, Western blot analysis (immunoblotting), ELISA, or fluorescence-activated cell sorting (FACS). Antibodies directed to HPR1 and/or HPR2 polypeptides can be prepared via

5 conventional antibody generation methods such as those described herein. Immunoprecipitation methods, Western blot (immunoblot) analysis, and enzyme-linked immunosorbent assays (ELISA) are standard in the art (see, for example, Ausubel, F. M. et al., Current Protocols in Molecular Biology, Volume 2, pp. 10.16.1-10.16.11, 10.8.1-10.8.21, and 11.2.1-11.2.22, John Wiley & Sons, Inc., 1991).

All publications and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference. Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it will be readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

## **Sequences Presented in the Sequence Listing**

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SEQ ID NO	Туре	Description
SEQ ID NO:1	Amino acid	Partial human HPR1 amino acid sequence
SEQ ID NO:2	Amino acid	Partial human HPR1 amino acid sequence
SEQ ID NO:3	Nucleotide	Human HPR1 cDNA sequence
SEQ ID NO:4	Amino acid	Human HPR1 amino acid sequence (745 amino acids)
SEQ ID NO:5	Nucleotide	Human HPR1 coding sequence
SEQ ID NO:6	Amino acid	Human LIF-R amino acid sequence (GenBank NP_002301)
SEQ ID NO:7	Amino acid	Human IL-12RB2 amino acid sequence (GenBank NP_001550)
SEQ ID NO:8	Amino acid	Human gp130 amino acid sequence (GenBank NP_002175)
SEQ ID NO:9	Amino acid	Human GCSFR amino acid sequence (SWISS-PROT Q99062)
SEQ ID NO:10	Amino acid	Portion of possible alternatively spliced form of human HPR1
SEQ ID NO:11	Amino acid	Portion of possible alternatively spliced form of human HPR1
SEQ ID NO:12	Amino acid	Mus musculus HPR1 amino acid sequence
SEQ ID NO:13	Amino acid	Possible 252-aa human HPR1 variant (WO 00/75314)
SEQ ID NO:14	Amino acid	Possible 652-aa human HPR1 variant (WO 00/75314)
SEQ ID NO:15	Amino acid	Possible 662-aa human HPR1 variant (WO 00/75314)
SEQ ID NO:16	Amino acid	Portion of possible alternatively spliced form of human HPR2
SEQ ID NO:17	Amino acid	Portion of possible alternatively spliced form of human HPR2
SEQ ID NO:18	Amino acid	Portion of possible alternatively spliced form of human HPR2
SEQ ID NO:19	Nucleotide	Human HPR2 cDNA sequence - exons 1 through 11
SEQ ID NO:20	Nucleotide	Human HPR2 coding sequence (encodes 629-aa form)
SEQ ID NO:21	Amino acid	Human HPR2 amino acid sequence (629 amino acids)
SEQ ID NO:22	Nucleotide	Human HPR2-ex8-ex9 coding sequence (encodes 565-aa form)
SEQ ID NO:23	Amino acid	Human HPR2-ex8-ex9 amino acid sequence (565 amino acids)
SEQ ID NO:24	Nucleotide	Human HPR2-ex9 coding sequence (encodes 356-aa form)
SEQ ID NO:25	Amino acid	Human HPR2-ex9 amino acid sequence (356 amino acids)
SEQ ID NO:26	Amino acid	Possible 384-aa human HPR2 variant (WO 00/73451)
SEQ ID NO:27	Amino acid	Mus musculus HPR2 amino acid sequence
SEQ ID NO:28	Nucleotide	Mus musculus HPR1 coding sequence
SEQ ID NO:29	Nucleotide	Mus musculus HPR2 coding sequence

5 <u>CLAIMS</u>

What is claimed is:

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1. An isolated polypeptide having HPR1 polypeptide activity comprising an amino acid sequence selected from the group consisting of:

- (a) the amino acid sequence of SEQ ID NO:4;
- 10 (b) an amino acid sequence selected from the group consisting of: amino acids 652 though 745 of SEQ ID NO:4, a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least 20 contiguous amino acids; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least 30 contiguous amino acids; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 that is at least 25% of the length of the sequence of amino acids 652 though 745 of SEQ ID NO:4; a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 that is at least 50% of the length of the sequence of amino acids 652 though 745 of SEQ ID NO:4; and a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4; and a fragment of the sequence of amino acids 652 though 745 of SEQ ID NO:4 comprising at least eight contiguous amino acids and comprising at least one tyrosine residue;
  - (c) an amino acid sequence comprising at least 8 amino acids and sharing amino acid identity with the amino acid sequences of (b), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99%, and at least 99.5%;
  - an amino acid sequence comprising both an amino acid sequence of (b) or (c), and an amino acid sequence selected from the group consisting of: amino acids 1 through 55 of SEQ ID NO:1; amino acids 56 through 77 of SEQ ID NO:1; amino acids 5 through 40 of SEQ ID NO:2; amino acids 1 through 32 of SEQ ID NO:4; amino acids 1 through 241 of SEQ ID NO:4; amino acids 1 through 525 of SEQ ID NO:4; amino acids 20 through 32 of SEQ ID NO:4; amino acids 33 through 134 of SEQ ID NO:4; amino acids Xaa1 through Xaa2 of SEQ ID NO:4, wherein Xaa1 is selected from the group consisting of amino acids 33 through 43 of SEQ ID NO:4 and Xaa2 is selected from the group consisting of amino acids 228 through 241 of SEQ ID NO:4; amino acids 33 through 238 of SEQ ID NO:4; amino acids 33 through 241 of SEQ ID NO:4; amino acids 33 through 525 of SEQ ID NO:4; amino acids 33 through 745 of SEQ ID NO:4; amino acids 44 through 94 of SEQ ID NO:4; amino acids 139 through 241 of SEQ ID NO:4; amino acids 242 through 326 of SEQ ID NO:4; amino acids 242 through 514 of SEQ ID NO:4; amino acids 337 through 419 of SEQ ID NO:4; amino acids 433 through 514 of SEQ ID NO:4; amino acids 526 through 556 of SEO ID NO:4; amino acids 533 through 552 of SEO ID NO:4; amino acids 553 through 745 of SEQ ID NO:4; amino acids 557 through 745 of SEQ ID NO:4; amino acids 563 through 573 of SEQ ID NO:4; amino acids 563 through 641 of SEQ ID NO:4; amino acids 567 through 581 of SEQ ID NO:4; amino acids 588 through 639 of SEQ ID NO:4; amino acids 631 through 641 of SEQ ID NO:4; SEQ ID NO:10; and SEQ ID NO:11;

5 (e) an amino acid sequence comprising both an amino acid sequence of (b) or (c), and a fragment of SEQ ID NO:4 comprising cytokine receptor domain amino acid sequences;

- (f) an allelic variant of any of (a)-(e); and
- (g) an amino acid sequence of (a)-(f), wherein a polypeptide comprising said amino acid sequence of (a)-(f) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (b)-(c).
  - 2. An isolated polypeptide having HPR1 polypeptide activity comprising an amino acid sequence selected from the group consisting of:
    - (a) the amino acid sequence of SEQ ID NO:12;

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- 15 (b) an amino acid sequence selected from the group consisting of: amino acids 633 though 726 of SEQ ID NO:12, a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 comprising at least 20 contiguous amino acids; a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 comprising at least 30 contiguous amino acids; a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 that is at least 25% of the length of the sequence of amino acids 633 though 726 of SEQ ID NO:12; a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 that is at least 50% of the length of the sequence of amino acids 633 though 726 of SEQ ID NO:12; and a fragment of the sequence of amino acids 633 though 726 of SEQ ID NO:12 comprising at least eight contiguous amino acids and comprising at least one tyrosine residue;
  - (c) an amino acid sequence comprising at least 8 amino acids and sharing amino acid identity with the amino acid sequences of (b), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99%, and at least 99.5%;
  - (d) an amino acid sequence comprising both an amino acid sequence of (b) or (c), and an amino acid sequence selected from the group consisting of: amino acids 1 through 28 of SEQ ID NO:12; amino acids 1 through 224 of SEQ ID NO:12; amino acids 1 through 509 of SEQ ID NO:12; amino acids 13 through 28 of SEQ ID NO:12; amino acids 29 through 124 of SEQ ID NO:12; amino acids Xaa1 through Xaa2 of SEQ ID NO:12, wherein Xaa1 is selected from the group consisting of amino acids 29 through 39 of SEQ ID NO:12 and Xaa2 is selected from the group consisting of amino acids 211 through 224 of SEQ ID NO:12; amino acids 29 through 128 of SEQ ID NO:12; amino acids 29 through 224 of SEQ ID NO:12; amino acids 29 through 509 of SEQ ID NO:12; amino acids 29 through 726 of SEQ ID NO:12; amino acids 225 through 499 of SEQ ID NO:12; amino acids 320 through 403 of SEQ ID NO:12; amino acids 417 through 508 of SEQ ID NO:12; amino acids 510 through 533 of SEQ ID NO:12; amino acids 517 through 532 of SEQ ID NO:12; amino acids 534 through 726 of SEQ ID NO:12; amino acids 547 through 557 of

5 SEQ ID NO:12; amino acids 547 through 622 of SEQ ID NO:12; and amino acids 612 through 622 of SEQ ID NO:12;

- (e) an amino acid sequence comprising both an amino acid sequence of (b) or (c), and a fragment of SEQ ID NO:12 comprising cytokine receptor domain amino acid sequences;
  - (f) an allelic variant of any of (a)-(e); and
- 10 (g) an amino acid sequence of (a)-(f), wherein a polypeptide comprising said amino acid sequence of (a)-(f) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (b)-(c).
- 3. An isolated polypeptide having HPR2 polypeptide activity comprising an amino acid sequence selected from the group consisting of:
  - (a) SEQ ID NO:23;
  - (b) SEQ ID NO:25;

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- (c) an amino acid sequence selected from the group consisting of: an amino acid sequence comprising at least 20 contiguous amino acids of SEQ ID NO:23 and comprising the contiguous amino acids 318 and 319 of SEQ ID NO:23; and amino acids 349 through 356 of SEQ ID NO:25;
  - (d) an amino acid sequence comprising at least 8 amino acids and sharing amino acid identity with the amino acid sequences of (c), wherein the percent amino acid identity is selected from the group consisting of: at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97.5%, at least 99%, and at least 99.5%;
  - (e) an amino acid sequence comprising both an amino acid sequence of (c) or (d), and an amino acid sequence selected from the group consisting of: amino acids 1 through 177 of SEQ ID NO:16; amino acids 216 through 245 of SEQ ID NO:16; SEQ ID NO:17; and SEQ ID NO:18;
  - (f) an amino acid sequence comprising both an amino acid sequence of (c) or (d), and an amino acid sequences of any of (a)-(b) comprising cytokine receptor domain amino acid sequences;
    - (g) an allelic variant of any of (a)-(f); and
- (h) an amino acid sequence of (a)-(g), wherein a polypeptide comprising said amino acid sequence of (a)-(g) binds to an antibody that also binds to a polypeptide comprising an amino acid sequence of any of (c)-(d).
- 4. An isolated polypeptide having HPR2 polypeptide activity comprising an amino acid sequence selected from the group consisting of:
  - (a) SEQ ID NO:27;
- 40 (b) SEQ ID NO:27 from which amino acids 297 through 316 or amino acids 317 through 336 have been deleted;

5 (c) an amino acid sequence comprising 20 or more contiguous amino acids of (a) or (b); and

- (d) an amino acid sequence comprising 30 or more contiguous amino and sharing at least 90% amino acid identity with the amino acid sequences of (a)-(b).
- An isolated nucleic acid encoding a polypeptide of any of claims 1 through 4.
  - 6. The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group consisting of:
    - (a) SEQ ID NO:3;
- 15 (b) SEQ ID NO:5;
  - (c) nucleotides 132 through 2366 of SEQ ID NO:3; and
  - (d) allelic variants of (a)-(c).
- 7. The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group consisting of nucleotides 1 through 137 of SEQ ID NO:3, nucleotides 138 through 228 of SEQ ID NO:3, nucleotides 229 through 346 of SEQ ID NO:3, nucleotides 347 through 528 of SEQ ID NO:3, nucleotides 529 through 680 of SEQ ID NO:3, nucleotides 681 through 846 of SEQ ID NO:3, nucleotides 847 through 926 of SEQ ID NO:3, nucleotides 927 through 1143 of SEQ ID NO:3, nucleotides 1144 through 1326 of SEQ ID NO:3, nucleotides 1327 through 1428 of SEQ ID NO:3, nucleotides 1429 through 1575 of SEQ ID NO:3, nucleotides 1576 through 1716 of SEQ ID NO:3, nucleotides 1717 through 1810 of SEQ ID NO:3, nucleotides 1811 through 1892 of SEQ ID NO:3, and nucleotides 1893 through 2480 of SEQ ID NO:3.
- 8. The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group 30 consisting of:
  - (a) SEQ ID NO:28; and
  - (b) allelic variants of (a).
- 9. The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group consisting of:
  - (a) SEQ ID NO:19;
  - (b) SEQ ID NO:20;
  - (c) SEQ ID NO:22;
  - (d) SEQ ID NO:24; and
- 40 (d) allelic variants of (a)-(d).

The nucleic acid of claim 5 comprising a nucleotide sequence selected from the group consisting of: nucleotides 107 through 175 of SEQ ID NO:19, nucleotides 107 through 478 of SEQ ID NO:19, nucleotides 107 through 1060 of SEQ ID NO:19, nucleotides 107 through 1099 of SEQ ID NO:19, nucleotides 107 through 1171 of SEQ ID NO:19, nucleotides 176 through 478 of SEQ ID NO:19, nucleotides 176 through 1099 of SEQ ID NO:19, nucleotides 176 through 1171 of SEQ ID NO:19, nucleotides 479 through 763 of SEQ ID NO:19, nucleotides 479 through 1099 of SEQ ID NO:19, nucleotides 503 through 1033 of SEQ ID NO:19, nucleotides 776 through 1066 of SEQ ID NO:19, nucleotides 776 through 1099 of SEQ ID NO:19, nucleotides 1232 through 1993 of SEQ ID NO:19, nucleotides 1283 through 1426 of SEQ ID NO:19, nucleotides 1283 through 1960 of SEQ ID NO:19, and nucleotides 1295 through 1939 of SEQ ID NO:19.

- 11. The nucleic acid of claim 5 comprising the nucleotide sequence of SEQ ID NO:29.
- 12. An isolated genomic nucleic acid corresponding to the nucleic acid of any of claims 5 to 11.
- 13. An expression vector comprising at least one nucleic acid according to any of claims 5 though 12.
- 14. A recombinant host cell comprising at least one nucleic acid according to any of claims 525 though 12.
  - 15. The recombinant host cell of claim 14, wherein the nucleic acid is integrated into the host cell genome.
- 30 16. A process for producing a polypeptide encoded by the nucleic acid of any of claims 5 though 12, comprising culturing a recombinant host cell under conditions promoting expression of said polypeptide, wherein the recombinant host cell comprises at least one nucleic acid according to any of claims 5 though 12.
- The process of claim 16 further comprising purifying said polypeptide.
  - 18. The polypeptide produced by the process of claim 17.

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- 19. An isolated antibody that binds to the polypeptide of any of claims 1 through 4 or claim 18.
- 20. The antibody of claim 19 wherein the antibody is a monoclonal antibody.

- 5 21. The antibody of claim 19 wherein the antibody is a human antibody.
  - 22. The antibody of claim 19 wherein the antibody is a humanized antibody.
- 23. The antibody of claim 19 wherein the antibody inhibits the activity of the polypeptide of any of claims 1 through 4 or claim 18.
- 24. A method of designing an inhibitor of the polypeptide of any of claims 1 through 4 or claim 18, the method comprising the steps of determining the three-dimensional structure of such polypeptide, analyzing the three-dimensional structure for the likely binding sites of substrates, synthesizing a molecule that incorporates a predicted reactive site, and determining the polypeptide-inhibiting activity of the molecule.
  - 25. A method for identifying compounds that alter HPR1 or HRP2 polypeptide activity comprising
- 20 (a) mixing a test compound with the polypeptide of any of claims 1 through 4 or claim 18; and
  - (b) determining whether the test compound alters the HPR1 or HRP2 polypeptide activity of said polypeptide.
- 26. A method for identifying compounds that inhibit the binding activity of HPR1 or HRP2 polypeptides comprising (a) mixing a test compound with the polypeptide of any of claims 1 through 4 or claim 18 and a binding partner of said polypeptide; and (b) determining whether the test compound inhibits the binding activity of said polypeptide.
- 30 27. A method for increasing ligand-binding activity comprising providing at least one compound selected from the group consisting of the polypeptide of any of claims 1 through 4 or claim 18 and agonists of said polypeptides.
- 28. The method of claim 27 wherein the method comprises increasing ligand-binding activity in a patient by administering to said patient at least one compound selected from the group consisting of the polypeptide of any of claims 1 through 4 or claim 18 and agonists of said polypeptides.
  - 29. A method for decreasing ligand-binding activity comprising providing at least one antagonist of the polypeptide of any of claims 1 through 4 or claim 18.

5 30. The method of claim 29 wherein the method comprises decreasing ligand-binding activity in a patient by administering at least one antagonist of the polypeptide of any of claims 1 through 4 or claim 18 to said patient.

- 31. The method of claim 29 wherein the antagonist is an antibody that inhibits the activity of said polypeptide.
  - 32. A method for treating a cell proliferation condition comprising administering at least one compound selected from the group consisting of the polypeptide of any of claims 1 through 4 or claim 18 and agonists of said polypeptides.
  - 33. The method of claim 32 wherein the cell proliferation condition is selected from the group consisting of pancytopenia, leukopenia, anemia, thrombocytopenia, neurodegenerative disorders, and osteoporosis resulting from a lack of bone-forming cells.

- 34. A method for treating a cell proliferation condition comprising administering an antagonist of the polypeptide of any of claims 1 through 4 or claim 18.
- 35. The method of claim 34 wherein the cell proliferation condition is selected from the group consisting of leukemia, tumour metastasis, and osteoporosis resulting from an excess of bone-resorbing cells.

#### SEQUENCE LISTING

<110> Immunex Corporation Cosman, David J. Mosley, Bruce A. Bird, Timothy A. DuBose, Robert F. Wiley, Steven R.

<120> HEMATOPOIETIN RECEPTORS HPR1 AND HPR2

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Pro 145	Val	Leu	Gly	Ile	Lys 150	Arg	Met	Ile	Gln	Ile 155	Glu	Trp	Ile	Lys	Pro 160
Glu	Leu	Ala	Pro	Val 165	Ser	Ser	Asp	Leu	Lys 170	Tyr	Thr	Leu	Arg	Phe 175	Arg
Thr	Val	Asn	Ser 180	Thr	Ser	Trp	Met	Glu 185	Val	Asn	Phe	Ala	Lys 190	Asn	Arg
Lys	Asp	Lys 195	Asn	Gln	Thr	Tyr	Asn 200	Leu	Thr	Gly	Leu	Gln 205	Pro	Phe	Thr
Glu	туr 210	Val	Ile	Ala	Leu	Arg 215	Cys	Ala	Val	Lys	Glu 220	Ser	Lys	Phe	Trp
Ser 225	Asp	Trp	Ser	Gln	Glu 230	Lys	Met	Gly	Met	Thr 235	Glu	Glu	Glu	Ala	Pro 240
Cys	Gly	Leu	Glu	Leu 245	Trp	Arg	Val	Leu	Lys 250	Pro	Ala	Glu	Ala	Asp 255	Gly
Arg	Arg	Pro	Val 260	Arg	Leu	Leu	Trp	Lys 265	Lys	Ala	Arg	Gly	Ala 270	Pro	Val
Leu	Glu	Lys 275	Thr	Leu	Gly	Tyr	Asn 280	Ile	Trp	Tyr	Tyr	Pro 285	Glu	Ser	Asn
Thr	Asn 290	Leu	Thr	Glu	Thr	Met 295	Asn	Thr	Thr	Asn	Gln 300	Gln	Leu	Glu	Leu
His 305	Leu	Gly	Gly	Glu	Ser 310	Phe	Trp	Val	Ser	Met 315	Ile	Ser	Tyr	Asn	Ser 320
Leu	Gly	Lys	Ser	Pro 325	Val	Ala	Thr	Leu	Arg 330	Ile	Pro	Ala	Ile	Gln 335	Glu

Lys	Ser	Phe	Gln 340	Cys	Ile	Glu	Val	Met 345	Gln	Ala	Cys	Val	Ala 350	Glu	Asp
Gln	Leu	Val 355	Val	Lys	Trp	Gln	Ser 360	Pro	Ala	Leu	Asp	Val 365	Asn	Thr	Trp
Met	Ile 370	Glu	Trp	Phe	Pro	Asp 375	Val	Asp	Ser	Glu	Pro 380	Thr	Thr	Leu	Ser
Trp 385	Glu	Ser	Val	Ser	Gln 390	Ala	Thr	Asn	Trp	Thr 395	Ile	Gln	Gln	Asp	Lys 400
Leu	Lys	Pro	Phe	Trp 405	Cys	Tyr	Asn	Ile	Ser 410	Val	Tyr	Pro	Met	Leu 415	His
Asp	Lys	Val	Gly 420	Glu	Pro	Tyr	Ser	Ile 425	Gln	Ala	Tyr	Ala	Lys 430	Glu	Gly
Val	Pro	Ser 435	Glu	Gly	Pro	Glu	Thr 440	Lys	Val	Glu	Asn	Ile 445	Gly	Val	Lys
Thr	Val 450	Thr	Ile	Thr	Trp	Lys 455	Glu	Ile	Pro	Lys	Ser 460	Glu	Arg	Lys	Gly
Ile 465	Ile	Cys	Asn	Tyr	Thr 470	Ile	Phe	Tyr	Gln	Ala 475	Glu	Gly	Gly	Lys	Gly 480
Phe	Ser	Lys	Thr	Val 485	Asn	Ser	Ser	Ile	Leu 490	Gln	Tyr	Gly	Leu	Glu 495	Ser
Leu	Lys	Arg	Lys 500	Thr	Ser	Tyr	Ile	Val 505	Gln	Val	Met	Ala	Ser 510	Thr	Ser
Ala	Gly	Gly 515	Thr	Asn	Gly	Thr	Ser 520	Ile	Asn	Phe	Lys	Thr 525	Ļeu	Ser	Phe
Ser	Val 530	Phe	Glu	Ile	Ile	Leu 535	Ile	Thr	Ser	Leu	Ile 540	Gly	Gly	Gly	Leu
Leu 545	Ile	Leu	Ile	Ile	Leu 550	Thr	Val	Ala	Tyr	Gly 555	Leu	Lys	Lys	Pro	Asn 560
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Gln	Glu	Asn	Asn	Leu 645	Gly	Gly	Glu	Lys	Asn 650	Gly	Tyr	Val	Thr	Cys 655	Pro	
Phe	Arg	Pro	Asp 660	Cys	Pro	Leu	Gly	Lys 665	Ser	Phe	Glu	Glu	Leu 670	Pro	Val	
Ser	Pro	Glu 675	Ile	Pro	Pro	Gly	Lys 680	Ser	Gln	Tyr	Leu	Arg 685	Ser	Arg	Met	
Pro	Glu 690	Gly	Thr	Arg	Pro	Glu 695	Ala	Lys	Glu	Gln	Leu 700	Leu	Phe	Ser	Gly	
Gln 705	Ser	Leu	Val	Pro	Asp 710	His	Leu	Cys	Glu	Glu 715	Gly	Ala	Pro	Asn	Pro 720	
Tyr	Leu	Lys	Asn	Ser 725	Val	Thr	Ala	Arg	Glu 730	Phe	Leu	Val	Ser	Glu 735	Lys	
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Phe Ile Leu Leu Tyr Leu Met Asn Gln Val Asn Ser Gln Lys Lys Gly 35 40 45

Ala Pro His Asp Leu Lys Cys Val Thr Asn Asn Leu Gln Val Trp Asn 50 55 60

Cys Ser Trp Lys Ala Pro Ser Gly Thr Gly Arg Gly Thr Asp Tyr Glu 65 70 75 80

Val Cys Ile Glu Asn Arg Ser Arg Ser Cys Tyr Gln Leu Glu Lys Thr 85 90 95

Ser Ile Lys Ile Pro Ala Leu Ser His Gly Asp Tyr Glu Ile Thr Ile 100 105 110

Asn Ser Leu His Asp Phe Gly Ser Ser Thr Ser Lys Phe Thr Leu Asn 115 120 125

Glu Gln Asn Val Ser Leu Ile Pro Asp Thr Pro Glu Ile Leu Asn Leu 130 135 140

Ser Ala Asp Phe Ser Thr Ser Thr Leu Tyr Leu Lys Trp Asn Asp Arg 145 150 155 160

Gly Ser Val Phe Pro His Arg Ser Asn Val Ile Trp Glu Ile Lys Val 165 170 175

Leu Arg Lys Glu Ser Met Glu Leu Val Lys Leu Val Thr His Asn Thr 180 185 190

Thr Leu Asn Gly Lys Asp Thr Leu His His Trp Ser Trp Ala Ser Asp

WO 02/029060	PCT/US01/31634

 Met
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 Asp Asp Leu His Phe 220
 Ser Gly Leu Glu Glu Trp 235
 Ser Asp Trp Ser Pro 240

Val Lys Asn Ile Ser Trp Ile Pro Asp Ser Gln Thr Lys Val Phe Pro 245 250 255

Gln Asp Lys Val Ile Leu Val Gly Ser Asp Ile Thr Phe Cys Cys Val 260 265 270

Ser Gln Glu Lys Val Leu Ser Ala Leu Ile Gly His Thr Asn Cys Pro 275 280 285

Leu Ile His Leu Asp Gly Glu Asn Val Ala Ile Lys Ile Arg Asn Ile 290 295 300

Ser Val Ser Ala Ser Ser Gly Thr Asn Val Val Phe Thr Thr Glu Asp 305 310 315 320

Asn Ile Phe Gly Thr Val Ile Phe Ala Gly Tyr Pro Pro Asp Thr Pro 325 330 335

Gln Gln Leu Asn Cys Glu Thr His Asp Leu Lys Glu Ile Ile Cys Ser 340 345 350

Trp Asn Pro Gly Arg Val Thr Ala Leu Val Gly Pro Arg Ala Thr Ser 355 360 365

Tyr Thr Leu Val Glu Ser Phe Ser Gly Lys Tyr Val Arg Leu Lys Arg 370 375 380

Ala Glu Ala Pro Thr Asn Glu Ser Tyr Gln Leu Leu Phe Gln Met Leu 385 390 395 400

Pro Asn Gln Glu Ile Tyr Asn Phe Thr Leu Asn Ala His Asn Pro Leu 405 410 415

Gly Arg Ser Gln Ser Thr Ile Leu Val Asn Ile Thr Glu Lys Val Tyr 420 425 430

Pro His Thr Pro Thr Ser Phe Lys Val Lys Asp Ile Asn Ser Thr Ala 435 440 445

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Leu 465	Cys	Glu	Ile	Glu	Ile 470	Lys	Lys	Ser	Asn	Ser 475	Val	Gln	Glu	Gln	Arg 480
Asn	Val	Thr	Ile	Lys 485	Gly	Val	Glu	Asn	Ser 490	Ser	Tyr	Leu	Val	Ala 495	Leu
Asp	Lys	Leu	Asn 500	Pro	Tyr	Thr	Leu	Туr 505	Thr	Phe	Arg	Ile	Arg 510	Cys	Ser
Thr	Glu	Thr 515	Phe	Trp	Lys	Trp	Ser 520	Lys	Trp	Ser	Asn	Lys 525	Lys	Gln	His
Leu	Thr 530	Thr	Glu	Ala	Ser	Pro 535	Ser	Lys	Gly	Pro	Asp 540	Thr	Trp	Arg	Glu
Trp 545	Ser	Ser	Asp	Gly	Lys 550	Asn	Leu	Ile	Ile	Tyr 555	Trp	Lys	Pro	Leu	Pro 560
Ile	Asn	Glu	Ala	Asn 565	Gly	Lys	Ile	Leu	Ser 570	Tyr	Asn	Val	Ser	Cys 575	Ser
Ser	Asp	Glu	Glu 580	Thr	Gln	Ser	Leu	Ser 585	Glu	Ile	Pro	Asp	Pro 590	Gln	His
Lys	Ala	Glu 595	Ile	Arg	Leu	Asp	Lys 600	Asn	Asp	Tyr	Ile	Ile 605	Ser	Val	Val
	Lys 610		Ser		_		Ser	Pro	Pro	Ser	Lys 620	Ile	Ala	Ser	Met
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Tyr	Val	Ile	Lys 660	Trp	Cys	Asn	Ser	Ser 665	Arg	Ser	Glu	Pro	Cys 670	Leu	Met
Asp	Trp	Arg 675	Lys	Val	Pro	Ser	Asn 680	Ser	Thr	Glu	Thr	Val 685	Ile	Glu	Ser

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Val Ala Glu Arg Pro Glu Asp Arg Ser Asp Ala Glu Pro Glu Asn His 930 935 940

Val Val Val Ser Tyr Cys Pro Pro Ile Ile Glu Glu Glu Ile Pro Asn 945 950 955 960

Pro Ala Ala Asp Glu Ala Gly Gly Thr Ala Gln Val Ile Tyr Ile Asp 965 970 975

Val Gln Ser Met Tyr Gln Pro Gln Ala Lys Pro Glu Glu Glu Glu Glu 980 985 990

Asn Asp Pro Val Gly Gly Ala Gly Tyr Lys Pro Gln Met His Leu Pro 995 1000 1005

Ile Asn Ser Thr Val Glu Asp Ile Ala Ala Glu Glu Asp Leu Asp 1010 1015 1020

Lys Thr Ala Gly Tyr Arg Pro Gln Ala Asn Val Asn Thr Trp Asn 1025 1030 1035

Leu Val Ser Pro Asp Ser Pro Arg Ser Ile Asp Ser Asn Ser Glu 1040 1045 1050

Ile Val Ser Phe Gly Ser Pro Cys Ser Ile Asn Ser Arg Gln Phe 1055 1060 1065

Leu Ile Pro Pro Lys Asp Glu Asp Ser Pro Lys Ser Asn Gly Gly 1070 1075 1080

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<213> Homo sapiens

<400> 7

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Thr Pro Val Ser Leu Lys Val Ser Thr Asn Ser Thr Arg Gln Ser Leu 35 40 45

His Leu Gln Trp Thr Val His Asn Leu Pro Tyr His Gln Glu Leu Lys Met Val Phe Gln Ile Gln Ile Ser Arg Ile Glu Thr Ser Asn Val Ile Trp Val Gly Asn Tyr Ser Thr Thr Val Lys Trp Asn Gln Val Leu His Trp Ser Trp Glu Ser Glu Leu Pro Leu Glu Cys Ala Thr His Phe Val Arg Ile Lys Ser Leu Val Asp Asp Ala Lys Phe Pro Glu Pro Asn Phe Trp Ser Asn Trp Ser Ser Trp Glu Glu Val Ser Val Gln Asp Ser Thr Gly Gln Asp Ile Leu Phe Val Phe Pro Lys Asp Lys Leu Val Glu Glu Gly Thr Asn Val Thr Ile Cys Tyr Val Ser Arg Asn Ile Gln Asn Asn Val Ser Cys Tyr Leu Glu Gly Lys Gln Ile His Gly Glu Gln Leu Asp Pro His Val Thr Ala Phe Asn Leu Asn Ser Val Pro Phe Ile Arg Asn Lys Gly Thr Asn Ile Tyr Cys Glu Ala Ser Gln Gly Asn Val Ser Glu Gly Met Lys Gly Ile Val Leu Phe Val Ser Lys Val Leu Glu Glu Pro Lys Asp Phe Ser Cys Glu Thr Glu Asp Phe Lys Thr Leu His Cys Thr Trp Asp Pro Gly Thr Asp Thr Ala Leu Gly Trp Ser Lys Gln Pro Ser Gln Ser Tyr Thr Leu Phe Glu Ser Phe Ser Gly Glu Lys Lys Leu Cys 

Thr His Lys Asn Trp Cys Asn Trp Gln Ile Thr Gln Asp Ser Gln Glu Thr Tyr Asn Phe Thr Leu Ile Ala Glu Asn Tyr Leu Arg Lys Arg Ser Val Asn Ile Leu Phe Asn Leu Thr His Arg Val Tyr Leu Met Asn Pro Phe Ser Val Asn Phe Glu Asn Val Asn Ala Thr Asn Ala Ile Met Thr Trp Lys Val His Ser Ile Arg Asn Asn Phe Thr Tyr Leu Cys Gln Ile Glu Leu His Gly Glu Gly Lys Met Met Gln Tyr Asn Val Ser Ile Lys Val Asn Gly Glu Tyr Phe Leu Ser Glu Leu Glu Pro Ala Thr Glu Tyr Met Ala Arg Val Arg Cys Ala Asp Ala Ser His Phe Trp Lys Trp Ser Glu Trp Ser Gly Gln Asn Phe Thr Thr Leu Glu Ala Pro Ser Glu Ala Pro Asp Val Trp Arg Ile Val Ser Leu Glu Pro Gly Asn His Thr Val Thr Leu Phe Trp Lys Pro Leu Ser Lys Leu His Ala Asn Gly Lys Ile Leu Phe Tyr Asn Val Val Glu Asn Leu Asp Lys Pro Ser Ser Ser Glu Leu His Ser Ile Pro Ala Pro Ala Asn Ser Thr Lys Leu Ile Leu Asp Arg Cys Ser Tyr Gln Ile Cys Val Ile Ala Asn Asn Ser Val Gly Ala Ser Pro Ala Ser Val Ile Val Ile Ser Ala Asp Pro Glu Asn 

Lys Glu Val Glu Glu Glu Arg Ile Ala Gly Thr Glu Gly Gly Phe Ser Leu Ser Trp Lys Pro Gln Pro Gly Asp Val Ile Gly Tyr Val Val Asp Trp Cys Asp His Thr Gln Asp Val Leu Gly Asp Phe Gln Trp Lys Asn Val Gly Pro Asn Thr Thr Ser Thr Val Ile Ser Thr Asp Ala Phe Arg Pro Gly Val Arg Tyr Asp Phe Arg Ile Tyr Gly Leu Ser Thr Lys Arg Ile Ala Cys Leu Leu Glu Lys Lys Thr Gly Tyr Ser Gln Glu Leu Ala Pro Ser Asp Asn Pro His Val Leu Val Asp Thr Leu Thr Ser His Ser Phe Thr Leu Ser Trp Lys Asp Tyr Ser Thr Glu Ser Gln Pro Gly Phe Ile Gln Gly Tyr His Val Tyr Leu Lys Ser Lys Ala Arg Gln Cys His Pro Arg Phe Glu Lys Ala Val Leu Ser Asp Gly Ser Glu Cys Cys Lys Tyr Lys Ile Asp Asn Pro Glu Glu Lys Ala Leu Ile Val Asp Asn Leu Lys Pro Glu Ser Phe Tyr Glu Phe Phe Ile Thr Pro Phe Thr Ser Ala Gly Glu Gly Pro Ser Ala Thr Phe Thr Lys Val Thr Thr Pro Asp Glu His Ser Ser Met Leu Ile His Ile Leu Leu Pro Met Val Phe Cys Val Leu Leu Ile Met Val Met Cys Tyr Leu Lys Ser Gln Trp Ile Lys Glu Thr Cys Tyr Pro Asp Ile Pro Asp Pro Tyr Lys Ser Ser Ile Leu Ser

770 775 780

Leu Ile Lys Phe Lys Glu Asn Pro His Leu Ile Ile Met Asn Val Ser 785 790 795 800

Asp Cys Ile Pro Asp Ala Ile Glu Val Val Ser Lys Pro Glu Gly Thr 805 810 815

Lys Ile Gln Phe Leu Gly Thr Arg Lys Ser Leu Thr Glu Thr Glu Leu 820 825 830

Thr Lys Pro Asn Tyr Leu Tyr Leu Pro Thr Glu Lys Asn His Ser 835 840 845

Gly Pro Gly Pro Cys Ile Cys Phe Glu Asn Leu Thr Tyr Asn Gln Ala 850 855 860

Ala Ser Asp Ser Gly Ser Cys Gly His Val Pro Val Ser Pro Lys Ala 865 870 875 880

Pro Ser Met Leu Gly Leu Met Thr Ser Pro Glu Asn Val Leu Lys Ala 885 890 895

Leu Glu Lys Asn Tyr Met Asn Ser Leu Gly Glu Ile Pro Ala Gly Glu 900 905 910

Thr Ser Leu Asn Tyr Val Ser Gln Leu Ala Ser Pro Met Phe Gly Asp 915 920 925

Lys Asp Ser Leu Pro Thr Asn Pro Val Glu Ala Pro His Cys Ser Glu 930 935 940

Tyr Lys Met Gln Met Ala Val Ser Leu Arg Leu Ala Leu Pro Pro 945 950 955 960

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His Tyr Cys

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<212> PRT

<213> Homo sapiens

<400> 8

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Lys Leu Thr Trp Thr Asn Pro Ser Ile Lys Ser Val Ile Ile Leu Lys Tyr Asn Ile Gln Tyr Arg Thr Lys Asp Ala Ser Thr Trp Ser Gln Ile Pro Pro Glu Asp Thr Ala Ser Thr Arg Ser Ser Phe Thr Val Gln Asp Leu Lys Pro Phe Thr Glu Tyr Val Phe Arg Ile Arg Cys Met Lys Glu Asp Gly Lys Gly Tyr Trp Ser Asp Trp Ser Glu Glu Ala Ser Gly Ile Thr Tyr Glu Asp Arg Pro Ser Lys Ala Pro Ser Phe Trp Tyr Lys Ile Asp Pro Ser His Thr Gln Gly Tyr Arg Thr Val Gln Leu Val Trp Lys Thr Leu Pro Pro Phe Glu Ala Asn Gly Lys Ile Leu Asp Tyr Glu Val Thr Leu Thr Arg Trp Lys Ser His Leu Gln Asn Tyr Thr Val Asn Ala Thr Lys Leu Thr Val Asn Leu Thr Asn Asp Arg Tyr Leu Ala Thr Leu Thr Val Arg Asn Leu Val Gly Lys Ser Asp Ala Ala Val Leu Thr Ile Pro Ala Cys Asp Phe Gln Ala Thr His Pro Val Met Asp Leu Lys Ala Phe Pro Lys Asp Asn Met Leu Trp Val Glu Trp Thr Thr Pro Arg Glu Ser Val Lys Lys Tyr Ile Leu Glu Trp Cys Val Leu Ser Asp Lys Ala Pro Cys Ile Thr Asp Trp Gln Glu Asp Gly Thr Val His Arg Thr Tyr Leu Arg Gly Asn Leu Ala Glu Ser Lys Cys Tyr Leu Ile Thr Val

PCT/US01/31634

485 490 495

WO 02/029060

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Tyr Leu Lys Gln Ala Pro Pro Ser Lys Gly Pro Thr Val Arg Thr Lys 515 520 525

Lys Val Gly Lys Asn Glu Ala Val Leu Glu Trp Asp Gln Leu Pro Val 530 540

Asp Val Gln Asn Gly Phe Ile Arg Asn Tyr Thr Ile Phe Tyr Arg Thr 545 550 555 560

Ile Ile Gly Asn Glu Thr Ala Val Asn Val Asp Ser Ser His Thr Glu 565 570 575

Tyr Thr Leu Ser Ser Leu Thr Ser Asp Thr Leu Tyr Met Val Arg Met 580 585 590

Ala Ala Tyr Thr Asp Glu Gly Gly Lys Asp Gly Pro Glu Phe Thr Phe 595 600 605

Thr Thr Pro Lys Phe Ala Gln Gly Glu Ile Glu Ala Ile Val Val Pro 610 615 620

Val Cys Leu Ala Phe Leu Leu Thr Thr Leu Leu Gly Val Leu Phe Cys 625 630 635 640

Phe Asn Lys Arg Asp Leu Ile Lys Lys His Ile Trp Pro Asn Val Pro 645 650 655

Asp Pro Ser Lys Ser His Ile Ala Gln Trp Ser Pro His Thr Pro Pro 660 665 670

Arg His Asn Phe Asn Ser Lys Asp Gln Met Tyr Ser Asp Gly Asn Phe 675 680 685

Thr Asp Val Ser Val Val Glu Ile Glu Ala Asn Asp Lys Lys Pro Phe 690 695 700

Pro Glu Asp Leu Lys Ser Leu Asp Leu Phe Lys Lys Glu Lys Ile Asn 705 710 715 720

Thr Glu Gly His Ser Ser Gly Ile Gly Gly Ser Ser Cys Met Ser Ser 725 730 735

Ser Arg Pro Ser Ile Ser Ser Ser Asp Glu Asn Glu Ser Ser Gln Asn 740 745 750

Thr Ser Ser Thr Val Gln Tyr Ser Thr Val Val His Ser Gly Tyr Arg
755 760 765

His Gln Val Pro Ser Val Gln Val Phe Ser Arg Ser Glu Ser Thr Gln 770 775 780

Pro Leu Leu Asp Ser Glu Glu Arg Pro Glu Asp Leu Gln Leu Val Asp 785 790 795 800

His Val Asp Gly Gly Asp Gly Ile Leu Pro Arg Gln Gln Tyr Phe Lys 805 810 815

Gln Asn Cys Ser Gln His Glu Ser Ser Pro Asp Ile Ser His Phe Glu 820 825 830

Arg Ser Lys Gln Val Ser Ser Val Asn Glu Glu Asp Phe Val Arg Leu 835 840 845

Lys Gln Gln Ile Ser Asp His Ile Ser Gln Ser Cys Gly Ser Gly Gln 850 855 860

Met Lys Met Phe Gln Glu Val Ser Ala Ala Asp Ala Phe Gly Pro Gly 865 870 875 880

Thr Glu Gly Gln Val Glu Arg Phe Glu Thr Val Gly Met Glu Ala Ala 885 890 895

Thr Asp Glu Gly Met Pro Lys Ser Tyr Leu Pro Gln Thr Val Arg Gln 900 905 910

Gly Gly Tyr Met Pro Gln 915

<210> 9

<211> 836

<212> PRT

<213> Homo sapiens

<400> 9

Met Ala Arg Leu Gly Asn Cys Ser Leu Thr Trp Ala Ala Leu Ile Ile 1 5 10 15

Leu Leu Pro Gly Ser Leu Glu Glu Cys Gly His Ile Ser Val Ser Ala Pro Ile Val His Leu Gly Asp Pro Ile Thr Ala Ser Cys Ile Ile Lys Gln Asn Cys Ser His Leu Asp Pro Glu Pro Gln Ile Leu Trp Arg Leu Gly Ala Glu Leu Gln Pro Gly Gly Arg Gln Gln Arg Leu Ser Asp Gly Thr Gln Glu Ser Ile Ile Thr Leu Pro His Leu Asn His Thr Gln Ala Phe Leu Ser Cys Cys Leu Asn Trp Gly Asn Ser Leu Gln Ile Leu Asp Gln Val Glu Leu Arg Ala Gly Tyr Pro Pro Ala Ile Pro His Asn Leu Ser Cys Leu Met Asn Leu Thr Thr Ser Ser Leu Ile Cys Gln Trp Glu Pro Gly Pro Glu Thr His Leu Pro Thr Ser Phe Thr Leu Lys Ser Phe Lys Ser Arg Gly Asn Cys Gln Thr Gln Gly Asp Ser Ile Leu Asp Cys Val Pro Lys Asp Gly Gln Ser His Cys Cys Ile Pro Arg Lys His Leu Leu Tyr Gln Asn Met Gly Ile Trp Val Gln Ala Glu Asn Ala Leu Gly Thr Ser Met Ser Pro Gln Leu Cys Leu Asp Pro Met Asp Val Val Lys Leu Glu Pro Pro Met Leu Arg Thr Met Asp Pro Ser Pro Glu Ala Ala Pro Pro Gln Ala Gly Cys Leu Gln Leu Cys Trp Glu Pro Trp 

Gln Pro Gly Leu His Ile Asn Gln Lys Cys Glu Leu Arg His Lys Pro

260 265 270

Gln Arg Gly Glu Ala Ser Trp Ala Leu Val Gly Pro Leu Pro Leu Glu 275 280 285

Ala Leu Gln Tyr Glu Leu Cys Gly Leu Leu Pro Ala Thr Ala Tyr Thr 290 295 300

Leu Gln Ile Arg Cys Ile Arg Trp Pro Leu Pro Gly His Trp Ser Asp 305 310 315 320

Trp Ser Pro Ser Leu Glu Leu Arg Thr Thr Glu Arg Ala Pro Thr Val 325 330 335

Arg Leu Asp Thr Trp Trp Arg Gln Arg Gln Leu Asp Pro Arg Thr Val $340 \hspace{1.5cm} 345 \hspace{1.5cm} 350$ 

Gln Leu Phe Trp Lys Pro Val Pro Leu Glu Glu Asp Ser Gly Arg Ile 355 360 365

Gln Gly Tyr Val Val Ser Trp Arg Pro Ser Gly Gln Ala Gly Ala Ile  $370 \hspace{1cm} 375 \hspace{1cm} 380$ 

Leu Pro Leu Cys Asn Thr Thr Glu Leu Ser Cys Thr Phe His Leu Pro 385 390 395 400

Ser Glu Ala Gln Glu Val Ala Leu Val Ala Tyr Asn Ser Ala Gly Thr 405 410 415

Ser Arg Pro Thr Pro Val Val Phe Ser Glu Ser Arg Gly Pro Ala Leu 420 425 430

Thr Arg Leu His Ala Met Ala Arg Asp Pro His Ser Leu Trp Val Gly 435 440 445

Trp Glu Pro Pro Asn Pro Trp Pro Gln Gly Tyr Val Ile Glu Trp Gly 450 455 460

Leu Gly Pro Pro Ser Ala Ser Asn Ser Asn Lys Thr Trp Arg Met Glu 465 470 475 480

Gln Asn Gly Arg Ala Thr Gly Phe Leu Leu Lys Glu Asn Ile Arg Pro 485 490 495

Phe Gln Leu Tyr Glu Ile Ile Val Thr Pro Leu Tyr Gln Asp Thr Met 500 505 510

Gly Pro Ser Gln His Val Tyr Ala Tyr Ser Gln Glu Met Ala Pro Ser His Ala Pro Glu Leu His Leu Lys His Ile Gly Lys Thr Trp Ala Gln Leu Glu Trp Val Pro Glu Pro Pro Glu Leu Gly Lys Ser Pro Leu Thr His Tyr Thr Ile Phe Trp Thr Asn Ala Gln Asn Gln Ser Phe Ser Ala Ile Leu Asn Ala Ser Ser Arg Gly Phe Val Leu His Gly Leu Glu Pro Ala Ser Leu Tyr His Ile His Leu Met Ala Ala Ser Gln Ala Gly Ala Thr Asn Ser Thr Val Leu Thr Leu Met Thr Leu Thr Pro Glu Gly Ser Glu Leu His Ile Ile Leu Gly Leu Phe Gly Leu Leu Leu Leu Thr Cys Leu Cys Gly Thr Ala Trp Leu Cys Cys Ser Pro Asn Arg Lys Asn Pro Leu Trp Pro Ser Val Pro Asp Pro Ala His Ser Ser Leu Gly Ser Trp Val Pro Thr Ile Met Glu Glu Asp Ala Phe Gln Leu Pro Gly Leu Gly Thr Pro Pro Ile Thr Lys Leu Thr Val Leu Glu Glu Asp Glu Lys Lys Pro Val Pro Trp Glu Ser His Asn Ser Ser Glu Thr Cys Gly Leu Pro Thr Leu Val Gln Thr Tyr Val Leu Gln Gly Asp Pro Arg Ala Val Ser Thr Gln Pro Gln Ser Gln Ser Gly Thr Ser Asp Gln Val Leu Tyr 

Gly Gln Leu Leu Gly Ser Pro Thr Ser Pro Gly Pro Gly His Tyr Leu 760 Arg Cys Asp Ser Thr Gln Pro Leu Leu Ala Gly Leu Thr Pro Ser Pro 775 Lys Ser Tyr Glu Asn Leu Trp Phe Gln Ala Ser Pro Leu Gly Thr Leu 790 795 Val Thr Pro Ala Pro Ser Gln Glu Asp Asp Cys Val Phe Gly Pro Leu 805 810 Leu Asn Phe Pro Leu Gln Gly Ile Arg Val His Gly Met Glu Ala 825 820 Leu Gly Ser Phe 835 <210> 10 <211> 7 <212> PRT <213> Homo sapiens <400> 10 Trp Lys Ser Thr Ser Val Lys 5 <210> 11 <211> 15 <212> PRT <213> Homo sapiens <400> 11 Glu Gly Lys Leu Peo Ala Ile Pro Val Leu Ser Ala Leu Lys 5 10 <210> 12 <211> 726 <212> PRT <213> Mus musculus <400> 12 Met Lys Pro Leu Gly Val Asn Ala Gly Ile Met Trp Thr Leu Ala Leu 5

Trp Ala Phe Ser Phe Leu Cys Lys Phe Ser Leu Ala Val Leu Pro Thr 20 25 30

Lys	Pro	Glu 35	Asn	Ile	Ser	Cys	Val 40	Phe	Tyr	Phe	Asp	Arg 45	Asn	Leu	Thr
Cys	Thr 50	Trp	Arg	Pro	Glu	Lys 55	Glu	Thr	Asn	Asp	Thr 60	Ser	Tyr	Ile	Val
Thr 65	Leu	Thr	Tyr	Ser	Tyr 70	Gly	Lys	Ser	Asn	Tyr 75	Ser	Asp	Asn	Ala	Thr 80
Glu	Ala	Ser	Tyr	Ser 85	Phe	Pro	Arg	Ser	Cys 90	Ala	Met	Pro	Pro	Asp 95	Ile
Cys	Ser	Val	Glu 100	Val	Gln	Ala	Gln	Asn 105	Gly	Asp	Gly	Lys	Val 110	Lys	Ser
Asp	Ile	Thr 115	Tyr	Trp	His	Leu	Ile 120	Ser	Ile	Ala	Lys	Thr 125	Glu	Pro	Pro
Ile	Ile 130	Leu	Ser	Val	Asn	Pro 135	Ile	Cys	Asn	Arg	Met 140	Phe	Gln	Ile	Gln
Trp 145	Lys	Pro	Arg	Glu	Lys 150	Thr	Arg	Gly	Phe	Pro 155	Leu	Val	Cys	Met	Leu 160
Arg	Phe	Arg	Thr	Val 165	Asn	Ser	Ser	Arg	Trp 170	Thr	Glu	Val	Asn	Phe 175	Glu
Asn	Cys	Lys	Gln 180	Val	Cys	Asn	Leu	Thr 185	Gly	Leu	Gln	Ala	Phe 190	Thr	Glu
Tyr	Val	Leu 195	Ala	Leu	Arg	Phe	Arg 200	Phe	Asn	Asp	Ser	Arg 205	Tyr	Trp	Ser
Lys	Trp 210	Ser	Lys	Glu	Glu	Thr 215	Arg	Val	Thr	Met	Glu 220	Glu	Val	Pro	His
Val 225	Leu	Asp	Leu	Trp	Arg 230	Ile	Leu	Glu	Pro	Ala 235	Asp	Met	Asn	Gly	Asp 240
Arg	Lys	Val	Arg	Leu 245	Leu	Trp	Lys	Lys	Ala 250	Arg	Gly	Ala	Pro	Val 255	Leu
Glu	Lys	Thr	Phe 260	Gly	Tyr	His	Ile	Gln 265	Tyr	Phe	Ala	Glu	Asn 270	Ser	Thr

Asn Leu Thr Glu Ile Asn Asn Ile Thr Thr Gln Gln Tyr Glu Leu Leu Leu Met Ser Gln Ala His Ser Val Ser Val Thr Ser Phe Asn Ser Leu Gly Lys Ser Gln Glu Thr Ile Leu Arg Ile Pro Asp Val His Glu Lys Thr Phe Gln Tyr Ile Lys Ser Met Gln Ala Tyr Ile Ala Glu Pro Leu Leu Val Val Asn Trp Gln Ser Ser Ile Pro Ala Val Asp Thr Trp Ile Val Glu Trp Leu Pro Glu Ala Ala Met Ser Lys Phe Pro Ala Leu Ser Trp Glu Ser Val Ser Gln Val Thr Asn Trp Thr Ile Glu Gln Asp Lys Leu Lys Pro Phe Thr Cys Tyr Asn Ile Ser Val Tyr Pro Val Leu Gly His Arg Val Gly Glu Pro Tyr Ser Ile Gln Ala Tyr Ala Lys Glu Gly Thr Pro Leu Lys Gly Pro Glu Thr Arg Val Glu Asn Ile Gly Leu Arg Thr Ala Thr Ile Thr Trp Lys Glu Ile Pro Lys Ser Ala Arg Asn Gly Phe Ile Asn Asn Tyr Thr Val Phe Tyr Gln Ala Glu Gly Gly Lys Glu Leu Ser Lys Thr Val Asn Ser His Ala Leu Gln Cys Asp Leu Glu Ser Leu Thr Arg Arg Thr Ser Tyr Thr Val Trp Val Met Ala Ser Thr Arg Ala Gly Gly Thr Asn Gly Val Arg Ile Asn Phe Lys Thr Leu Ser Ile Ser Val Phe Glu Val Val Leu Leu Thr Ser Leu Val Gly Gly Leu

515 520 525

Leu Leu Ser Ile Lys Thr Val Thr Phe Gly Leu Arg Lys Pro Asn 530 535 540

Arg Leu Thr Pro Leu Cys Cys Pro Asp Val Pro Asn Pro Ala Glu Ser 545 550 555 560

Ser Leu Ala Thr Trp Leu Gly Asp Gly Phe Lys Lys Ser Asn Met Lys 565 570 575

Glu Thr Gly Asn Ser Gly Asn Thr Glu Asp Val Val Leu Lys Pro Cys 580 585 590

Pro Val Pro Ala Asp Leu Ile Asp Lys Leu Val Val Asn Phe Glu Asn 595 600 605

Phe Leu Glu Val Val Leu Thr Glu Glu Ala Gly Lys Gly Gln Ala Ser 610 615 620

Ile Leu Gly Gly Glu Ala Asn Glu Tyr Val Thr Ser Pro Ser Arg Pro 625 630 635 640

Asp Gly Pro Pro Gly Lys Ser Phe Lys Glu Pro Ser Ile Leu Thr Glu 645 650 655

Val Ala Ser Glu Asp Ser His Ser Thr Cys Ser Arg Met Ala Asp Glu 660 665 670

Ala Tyr Ser Glu Leu Ala Arg Gln Pro Ser Ser Ser Cys Gln Ser Pro 675 680 685

Gly Leu Ser Pro Pro Arg Glu Asp Gln Ala Gln Asn Pro Tyr Leu Lys 690 695 700

Asn Ser Val Thr Thr Arg Glu Phe Leu Val His Glu Asn Ile Pro Glu 705 710 715 720

His Ser Lys Gly Glu Val 725

<210> 13

<211> 252

<212> PRT

<213> Homo sapiens

<400> 13

Met 1	Lys	Leu	Ser	Pro 5	Gln	Pro	Ser	Cys	Val 10	Asn	Leu	Gly	Met	Met 15	Trp
Thr	Trp	Ala	Leu 20	Trp	Met	Leu	Pro	Ser 25	Leu	Cys	Lys	Phe	Ser 30	Leu	Ala
Ala	Leu	Pro 35	Ala	Lys	Pro	Glu	Asn 40	Ile	Ser	Cys	Val	Tyr 45	Tyr	Tyr	Arg
Lys	Asn 50	Leu	Thr	Cys	Thr	Trp 55	Ser	Pro	Gly	Lys	Glu 60	Thr	Ser	Tyr	Thr
Gln 65	Tyr	Thr	Val	Lys	Arg 70	Thr	Tyr	Ala	Phe	Gly 75	Glu	Lys	His	Asp	Asn 80
Cys	Thr	Thr	Asn	Ser 85	Ser	Thr	Ser	Glu	Asn 90	Arg	Ala	Ser	Cys	Ser 95	Phe
Phe	Leu	Pro	Arg 100	Ile	Thr	Ile	Pro	Asp 105	Asn	Tyr	Thr	Ile	Glu 110	Val	Glu
Ala	Glu	Asn 115	Gly	Asp	Gly	Val	Ile 120	Lys	Ser	His	Met	Thr 125	Tyr	Trp	Arg
Leu	Glu 130	Asn	Ile	Ala	Lys	Thr 135	Glu	Pro	Pro	Lys	Ile 140	Phe	Arg	Val	Lys
Pro 145	Val	Leu	Gly	Ile	Lys 150	Arg	Met	Ile	Gln	Ile 155	Glu	Trp	Ile	Lys	Pro 160
Glu	Leu	Ala	Pro	Val 165	Ser	Ser	Asp	Leu	Lys 170	Tyr	Thr	Leu	Arg	Phe 175	Arg
Thr	Val	Asn	Ser 180	Thr	Ser	Trp	Met	Glu 185	Val	Asn	Phe	Ala	Lys 190	Asn	Arg
Lys	Asp	Lys 195	Asn	Gln	Thr	Tyr	Asn 200	Leu	Thr	Gly	Leu	Gln 205	Pro	Phe	Thr
Glu	Tyr 210	Val	Ile	Ala	Leu	Arg 215	Cys	Ala	Val	Lys	Glu 220	Ser	Lys	Phe	Trp
Ser 225	Asp	Trp	Ser	Gln	Glu 230	Lys	Met	Gly	Met	Thr 235	Glu	Glu	Glu	Gly	Lys 240

Leu Leu Pro Ala Ile Pro Val Leu Ser Thr Leu Val 245 250

<210> 14

<211> 652

<212> PRT

<213> Homo sapiens

<400> 14

Met Lys Leu Ser Pro Gln Pro Ser Cys Val Asn Leu Gly Met Met Trp 1 5 10 15

Thr Trp Ala Leu Trp Met Leu Pro Ser Leu Cys Lys Phe Ser Leu Ala 20 25 30

Ala Leu Pro Ala Lys Pro Glu Asn Ile Ser Cys Val Tyr Tyr Tyr Arg  $35 \hspace{1cm} 40 \hspace{1cm} 45$ 

Lys Asn Leu Thr Cys Thr Trp Ser Pro Gly Lys Glu Thr Ser Tyr Thr 50 55 60

Gln Tyr Thr Val Lys Arg Thr Tyr Ala Phe Gly Glu Lys His Asp Asn 65 70 75 80

Cys Thr Thr Asn Ser Ser Thr Ser Glu Asn Arg Ala Ser Cys Ser Phe 85 90 95

Phe Leu Pro Arg Ile Thr Ile Pro Asp Asn Tyr Thr Ile Glu Val Glu
100 105 110

Ala Glu Asn Gly Asp Gly Val Ile Lys Ser His Met Thr Tyr Trp Arg 115 120 125

Leu Glu Asn Ile Ala Lys Thr Glu Pro Pro Lys Ile Phe Arg Val Lys 130 135 140

Pro Val Leu Gly Ile Lys Arg Met Ile Gln Ile Glu Trp Ile Lys Pro 145 150 155 160

Glu Leu Ala Pro Val Ser Ser Asp Leu Lys Tyr Thr Leu Arg Phe Arg 165 170 175

Thr Val Asn Ser Thr Ser Trp Met Glu Val Asn Phe Ala Lys Asn Arg 180 185 190

Lys Asp Lys Asn Gln Thr Tyr Asn Leu Thr Gly Leu Gln Pro Phe Thr 195 200 205

Glu	Tyr 210	Val	Ile	Ala	Leu	Arg 215	Cys	Ala	Val	Lys	Glu 220	Ser	Lys	Phe	Trp
Ser 225	Asp	Trp	Ser	Gln	Glu 230	Lys	Met	Gly	Met	Thr 235	Glu	Glu	Glu	Ala	Pro 240
Cys	Gly	Leu	Glu	Leu 245	Trp	Arg	Val	Leu	Lys 250	Pro	Ala	Glu	Ala	Asp 255	Gly
Arg	Arg	Pro	Val 260	Arg	Leu	Leu	Trp	Lys 265	Lys	Ala	Arg	Gly	Ala 270	Pro	Val
Leu	Glu	Lys 275	Thr	Leu	Gly	Tyr	Asn 280	Ile	Trp	Tyr	Tyr	Pro 285	Glu	Ser	Asn
Thr	Asn 290	Leu	Thr	Glu	Thr	Met 295	Asn	Thr	Thr	Asn	Gln 300	Gln	Leu	Glu	Leu
His 305	Leu	Gly	Gly	Glu	Ser 310	Phe	Trp	Val	Ser	Met 315	Ile	Ser	Tyr	Asn	Ser 320
Leu	Gly	Lys	Ser	Pro 325	Val	Ala	Thr	Leu	Arg 330	Ile	Pro	Ala	Ile	Gln 335	Glu
Lys	Ser	Phe	Gln 340	Cys	Ile	Glu	Val	Met 345	Gln	Ala	Cys	Val	Ala 350	Glu	Asp
Gln	Leu	Val 355	Val	Lys	Trp	Gln	Ser 360	Ser	Ala	Leu	Asp	Val 365	Asn	Thr	Trp
Met	Ile 370	Glu	Trp	Phe	Pro	Asp 375	Val	Asp	Ser	Glu	Pro 380	Thr	Thr	Leu	Ser
Trp 385	Glu	Ser	Val	Ser	Gln 390	Ala	Thr	Asn	Trp	Thr 395	Ile	Gln	Gln	Asp	Lys 400
Leu	Lys	Pro	Phe	Trp 405	Cys	Tyr	Asn	Ile	Ser 410	Val	Tyr	Pro	Met	Leu 415	His
Asp	Lys	Val	Gly 420	Glu	Pro	Tyr	Ser	Ile 425	Gln	Ala	Tyr	Ala	Lys 430	Glu	Gly
Val	Pro	Ser 435	Glu	Gly	Pro	Glu	Thr 440	Lys	Val	Glu	Asn	Ile 445	Gly	Val	Lys

Thr Val Thr Ile Thr Trp Lys Glu Ile Pro Lys Ser Glu Arg Lys Gly

Ile Ile Cys Asn Tyr Thr Ile Phe Tyr Gln Ala Glu Gly Gly Lys Gly 470 Phe Ser Lys Thr Val Asn Ser Ser Ile Leu Gln Tyr Gly Leu Glu Ser 485 490 Leu Lys Arg Lys Thr Ser Tyr Ile Val Gln Val Met Ala Asn Thr Ser 500 505 Ala Gly Gly Thr Asn Gly Thr Ser Ile Asn Phe Lys Thr Leu Ser Phe 520 525 Ser Val Phe Glu Ile Ile Leu Ile Thr Ser Leu Ile Gly Gly Leu Leu Ile Leu Ile Leu Thr Val Ala Tyr Gly Leu Lys Lys Pro Asn 545 550 555 Lys Leu Thr His Leu Cys Trp Pro Thr Val Pro Asn Pro Ala Glu Ser 565 570 Ser Ile Ala Thr Trp His Gly Asp Asp Phe Lys Asp Lys Leu Asn Leu 580 585 590 Lys Glu Ser Asp Asp Ser Val Asn Thr Glu Asp Arg Ile Leu Lys Pro 600 605 595 Cys Ser Thr Pro Ser Asp Lys Leu Val Ile Asp Lys Leu Val Val Asn 610 615 620 Phe Gly Asn Val Leu Gln Glu Ile Phe Thr Asp Glu Ala Arg Thr Gly 625 630 635 640 Gln Glu Lys Gln Phe Arg Arg Gly Lys Glu Trp Asp 645

<210> 15

<211> 662

<212> PRT

<213> Homo sapiens

<400> 15

Met Lys Leu Ser Pro Gln Pro Ser Cys Val Asn Leu Gly Met Met Trp

Thr Trp Ala Leu Trp Met Leu Pro Ser Leu Cys Lys Phe Ser Leu Ala Ala Leu Pro Ala Lys Pro Glu Asn Ile Ser Cys Val Tyr Tyr Arg Lys Asn Leu Thr Cys Thr Trp Ser Pro Gly Lys Glu Thr Ser Tyr Thr Gln Tyr Thr Val Lys Arg Thr Tyr Ala Phe Gly Glu Lys His Asp Asn Cys Thr Thr Asn Ser Ser Thr Ser Glu Asn Arg Ala Ser Cys Ser Phe Phe Leu Pro Arg Ile Thr Ile Pro Asp Asn Tyr Thr Ile Glu Val Glu Ala Glu Asn Gly Asp Gly Val Ile Lys Ser His Met Thr Tyr Trp Arg Leu Glu Asn Ile Ala Lys Thr Glu Pro Pro Lys Ile Phe Arg Val Lys Pro Val Leu Gly Ile Lys Arg Met Ile Gln Ile Glu Trp Ile Lys Pro Glu Leu Ala Pro Val Ser Ser Asp Leu Lys Tyr Thr Leu Arg Phe Arg Thr Val Asn Ser Thr Ser Trp Met Glu Val Asn Phe Ala Lys Asn Arg Lys Asp Lys Asn Gln Thr Tyr Asn Leu Thr Gly Leu Gln Pro Phe Thr Glu Tyr Val Ile Ala Leu Arg Cys Ala Val Lys Glu Ser Lys Phe Trp Ser Asp Trp Ser Gln Glu Lys Met Gly Met Thr Glu Glu Glu Ala Pro 

Cys Gly Leu Glu Leu Trp Arg Val Leu Lys Pro Ala Glu Ala Asp Gly

Arg Arg Pro Val Arg Leu Leu Trp Lys Lys Ala Arg Gly Ala Pro Val Leu Glu Lys Thr Leu Gly Tyr Asn Ile Trp Tyr Tyr Pro Glu Ser Asn Thr Asn Leu Thr Glu Thr Met Asn Thr Thr Asn Gln Gln Leu Glu Leu His Leu Gly Gly Glu Ser Phe Trp Val Ser Met Ile Ser Tyr Asn Ser Leu Gly Lys Ser Pro Val Ala Thr Leu Arg Ile Pro Ala Ile Gln Glu Lys Ser Phe Gln Cys Ile Glu Val Met Gln Ala Cys Val Ala Glu Asp Gln Leu Val Val Lys Trp Gln Ser Ser Ala Leu Asp Val Asn Thr Trp Met Ile Glu Trp Phe Pro Asp Val Asp Ser Glu Pro Thr Thr Leu Ser Trp Glu Ser Val Ser Gln Ala Thr Asn Trp Thr Ile Gln Gln Asp Lys Leu Lys Pro Phe Trp Cys Tyr Asn Ile Ser Val Tyr Pro Met Leu His Asp Lys Val Gly Glu Pro Tyr Ser Ile Gln Ala Tyr Ala Lys Glu Gly Val Pro Ser Glu Gly Pro Glu Thr Lys Val Glu Asn Ile Gly Val Lys Thr Val Thr Ile Thr Trp Lys Glu Ile Pro Lys Ser Glu Arg Lys Gly Ile Ile Cys Asn Tyr Thr Ile Phe Tyr Gln Ala Glu Gly Gly Lys Gly Phe Ser Lys Thr Val Asn Ser Ser Ile Leu Gln Tyr Gly Leu Glu Ser 

Leu Lys Arg Lys Thr Ser Tyr Ile Val Gln Val Met Ala Ser Thr Ser 500 505 510

Ala Gly Gly Thr Asn Gly Thr Ser Ile Asn Phe Lys Thr Leu Ser Phe 515 520 525

Ser Val Phe Glu Ile Ile Leu Ile Thr Ser Leu Ile Gly Gly Leu 530 535 540

Leu Ile Leu Ile Ile Leu Thr Val Ala Tyr Gly Leu Lys Lys Pro Asn 545 550 555 560

Lys Leu Thr His Leu Cys Trp Pro Thr Val Pro Asn Pro Ala Glu Ser 565 570 575

Ser Ile Ala Thr Trp His Gly Asp Asp Phe Lys Asp Lys Leu Asn Leu 580 585 590

Lys Glu Ser Asp Asp Ser Val Asn Thr Glu Asp Arg Ile Leu Lys Pro 595 600 605

Cys Ser Thr Pro Ser Asp Lys Leu Val Ile Asp Lys Leu Val Val Asn 610 615 620

Phe Gly Asn Val Leu Gln Glu Ile Phe Thr Asp Glu Ala Arg Thr Gly 625 630 635 640

Gln Glu Asn Asn Leu Gly Gly Glu Lys Asn Gly Thr Arg Ile Leu Ser 645 650 655

Ser Cys Pro Thr Ser Ile 660

<210> 16

<211> 344

<212> PRT

<213> Homo sapiens

<400> 16

Asn Pro Lys Asn Glu Ser Ser Glu Asn Ile Arg Glu Arg Leu Ser Leu 1 5 10 15

Pro Ser Thr Leu Gln Gln Asn Phe Gly Thr Leu Asn Phe Trp Phe Gln 20 25 30

Arg Ser His Asn Phe His Asn Leu Thr Thr Glu Glu Gly Pro Ser Thr

35 40 45

Pro Ile Gly Thr Leu Lys Pro Gly Leu Val Ile Lys Ala Val Arg Lys 50 60

Leu Leu Met Asn Asp Ser Asp Gln Gly Gly Lys Leu Thr Thr Gly Val 65 70 75 80

Phe Thr Pro Gln Gln Leu Ala Asn Thr Thr Asn Gln Gly Leu Ser Arg 85 90 95

Cys Leu Ser Arg Phe Lys Lys Val Ile Arg Ala Met Leu Met Met Lys
100 105 110

Ile Lys Leu Lys Arg Ile Thr Asn Ile Asn Cys Ser Gly His Ile Trp
115 120 125

Val Glu Pro Ala Thr Ile Phe Lys Met Gly Met Asn Ile Ser Ile Tyr 130 135 140

Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu His Phe Tyr 145 150 155 160

Lys Asn Gly Ile Lys Glu Arg Phe Gln Ile Thr Arg Ile Asn Lys Thr 165 170 175

Thr Ala Arg Leu Trp Tyr Lys Asn Phe Leu Glu Pro His Ala Ser Met 180 185 190

Tyr Cys Thr Ala Glu Cys Pro Lys His Phe Gln Glu Thr Leu Ile Cys 195 200 205

Gly Lys Asp Ile Ser Ser Gly Phe Cys Ile Thr Asp Tyr Ser Gln Lys 210 215 220

Pro Ser Gln Val Leu Ala Gly Gly Pro Leu Ser Pro Asn Pro Thr Pro 225 230 235 240

Gly Asn Val Glu Asp Pro Pro Asp Ile Pro Asp Glu Val Thr Cys Val
245 250 255

Ile Tyr Glu Tyr Ser Gly Asn Met Thr Cys Thr Trp Asn Ala Gly Lys 260 265 270

Leu Thr Tyr Ile Asp Thr Lys Tyr Val Val His Val Lys Ser Leu Glu 275 280 285

Thr Glu Glu Glu Gln Gln Tyr Leu Thr Ser Ser Tyr Ile Asn Ile Ser 290 295 Thr Asp Ser Leu Gln Gly Gly Lys Lys Tyr Leu Val Trp Val Gln Ala 305 310 315 Ala Asn Ala Leu Gly Met Glu Glu Ser Lys Gln Leu Gln Ile His Leu 330 325 Asp Asp Ile Ala Pro His Glu Arg 340 <210> 17 <211> 39 <212> PRT <213> Homo sapiens <400> 17 Ile Glu Asp Leu Ser Ile Asn Val Met Ala Ala Asn Ile Leu Glu Thr 10 Asn Asn Phe Leu Thr Arg Asp Thr Asn Met Lys Gln Ser Ala Phe Glu 25 30 20 Ser Gln Ile Phe Gly Thr Val 35 <210> 18 <211> 12 <212> PRT <213> Homo sapiens <400> 18 Ser Asn Trp Leu Ala Leu Lys Gly Asp Glu Glu Lys 5 <210> 19 <211> 2830 <212> DNA <213> Homo sapiens <400> 19 60 aaagaagaca tgacacagcc aacaagggtg gcagcctggc tctgaagtgg aattatgtgc ttcaaacagg ttgaaagagg gaaacagtct tttcctgctt ccagacatga atcaggtcac 120 tattcaatgg gatgcagtaa tagcccttta catactcttc agctggtgtc atggaggaat 180 240 tacaaatata aactgctctg gccacatctg ggtagaacca gccacaattt ttaagatggg

tatgaatatc	tctatatatt	gccaagcagc	aattaagaac	tgccaaccaa	ggaaacttca	300
tttttataaa	aatggcatca	aagaaagatt	tcaaatcaca	aggattaata	aaacaacagc	360
tcggctttgg	tataaaaact	ttctggaacc	acatgcttct	atgtactgca	ctgctgaatg	420
tcccaaacat	tttcaagaga	cactgatatg	tggaaaagac	atttcttctg	gatatccgcc	480
agatattcct	gatgaagtaa	cctgtgtcat	ttatgaatat	tcaggcaaca	tgacttgcac	540
ctggaatgct	gggaagctca	cctacataga	cacaaaatac	gtggtacatg	tgaagagttt	600
agagacagaa	gaagagcaac	agtatctcac	ctcaagctat	attaacatct	ccactgattc	660
attacaaggt	ggcaagaagt	acttggtttg	ggtccaagca	gcaaacgcac	taggcatgga	720
agagtcaaaa	caactgcaaa	ttcacctgga	tgatatagtg	ataccttctg	cagccgtcat	780
ttccagggct	gagactataa	atgctacagt	gcccaagacc	ataatttatt	gggatagtca	840
aacaacaatt	gaaaaggttt	cctgtgaaat	gagatacaag	gctacaacaa	accaaacttg	900
gaatgttaaa	gaatttgaca	ccaattttac	atatgtgcaa	cagtcagaat	tctacttgga	960
gccaaacatt	aagtacgtat	ttcaagtgag	atgtcaagaa	acaggcaaaa	ggtactggca	1020
gccttggagt	tcactgtttt	ttcataaaac	acctgaaaca	gttccccagg	tcacatcaaa	1080
agcattccaa	catgacacat	ggaattctgg	gctaacagtt	gcttccatct	ctacagggca	1140
ccttacttct	gacaacagag	gagacattgg	acttttattg	ggaatgatcg	tctttgctgt	1200
tatgttgtca	attctttctt	tgattgggat	atttaacaga	tcattccgaa	ctgggattaa	1260
aagaaggatc	ttattgttaa	taccaaagtg	gctttatgaa	gatattccta	atatgaaaaa	1320
cagcaatgtt	gtgaaaatgc	tacaggaaaa	tagtgaactt	atgaataata	attccagtga	1380
gcaggtccta	tatgttgatc	ccatgattac	agagataaaa	gaaatcttca	tcccagaaca	1440
caagcctaca	gactacaaga	aggagaatac	aggacccctg	gagacaagag	actacccgca	1500
aaactcgcta	ttcgacaata	ctacagttgt	atatattcct	gatctcaaca	ctggatataa	1560
accccaaatt	tcaaattttc	tgcctgaggg	aagccatctc	agcaataata	atgaaattac	1620
ttccttaaca	cttaaaccac	cagttgattc	cttagactca	ggaaataatc	ccaggttaca	1680
aaagcatcct	aattttgctt	tttctgtttc	aagtgtgaat	tcactaagca	acacaatatt	1740
tcttggagaa	ttaagcctca	tattaaatca	aggagaatgc	agttctcctg	acatacaaaa	1800
ctcagtagag	gaggaaacca	ccatgctttt	ggaaaatgat	tcacccagtg	aaactattcc	1860
agaacagacc	ctgcttcctg	atgaatttgt	ctcctgtttg	gggatcgtga	atgaggagtt	1920
gccatctatt	aatacttatt	ttccacaaaa	tattttggaa	agccacttca	ataggatttc	1980
actcttggaa	aagtagagct	gtgtggtcaa	aatcaatatg	agaaagctgc	cttgcaatct	2040

gaacttgggt	tttccctgca	atagaaattg	aattctgcct	ctttttgaaa	aaaatgtatt	2100
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Leu Phe Ser Trp Cys His Gly Gly Ile Thr Asn Ile Asn Cys Ser Gly 20 25 30

His Ile Trp Val Glu Pro Ala Thr Ile Phe Lys Met Gly Met Asn Ile 35 40 45

Ser Ile Tyr Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu 50 60

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<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Homo sapiens

<sup>&</sup>lt;400> 21

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Ala	Ser	Met	Tyr 100	Cys	Thr	Ala	Glu	Cys 105	Pro	Lys	His	Phe	Gln 110	Glu	Thr
Leu	Ile	Cys 115	Gly	Lys	Asp	Ile	Ser 120	Ser	Gly	Tyr	Pro	Pro 125	Asp	Ile	Pro
Asp	Glu 130	Val	Thr	Cys	Val	Ile 135	Tyr	Glu	Tyr	Ser	Gly 140	Asn	Met	Thr	Cys
Thr 145	Trp	Asn	Ala	Gly	Lys 150	Leu	Thr	Tyr	Ile	Asp 155	Thr	Lys	Tyr	Val	Val 160
His	Val	Lys	Ser	Leu 165	Glu	Thr	Glu	Glu	Glu 170	Gln	Gln	Tyr	Leu	Thr 175	Ser
Ser	Tyr	Ile	Asn 180	Ile	Ser	Thr	Asp	Ser 185	Leu	Gln	Gly	Gly	Lys 190	Lys	Tyr
Leu	Val	Trp 195	Val	Gln	Ala	Ala	Asn 200	Ala	Leu	Gly	Met	Glu 205	Glu	Ser	Lys
Gln	Leu 210	Gln	Ile	His	Leu	Asp 215	Asp	Ile	Val	Ile	Pro 220	Ser	Ala	Ala	Val
Ile 225	Ser	Arg	Ala	Glu	Thr 230	Ile	Asn	Ala	Thr	Val 235	Pro	Lys	Thr	Ile	Ile 240
Tyr	Trp	Asp	Ser	Gln 245	Thr	Thr	Ile	Glu	Lys 250	Val	Ser	Cys	Glu	Met 255	Arg
Tyr	Lys	Ala	Thr 260	Thr	Asn	Gln	Thr	Trp 265	Asn	Val	Lys	Glu	Phe 270	Asp	Thr
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Lys	Tyr 290	Val	Phe	Gln	Val	Arg 295	Cys	Gln	Glu	Thr	Gly 300	Lys	Arg	Tyr	Trp

Gln 305	Pro	Trp	Ser	Ser	10 310	Phe	Phe	His	Lys	Thr 315	Pro	Glu	Thr	Val	9ro 320
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His	Leu	Ser	Asn 500	Asn	Asn	Glu	Ile	Thr 505	Ser	Leu	Thr	Leu	Lys 510	Pro	Pro
Val	Asp	Ser 515	Leu	Asp	Ser	Gly	Asn 520	Asn	Pro	Arg	Leu	Gln 525	Lys	His	Pro
Asn	Phe 530	Ala	Phe	Ser	Val	Ser 535	Ser	Val	Asn	Ser	Leu 540	Ser	Asn	Thr	Ile
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Asn Asp Ser Pro Ser Glu Thr Ile Pro Glu Gln Thr Leu Leu Pro Asp 580 585 590

Glu Phe Val Ser Cys Leu Gly Ile Val Asn Glu Glu Leu Pro Ser Ile 595 600 605

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<211> 1698

<212> DNA

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Leu Phe Ser Trp Cys His Gly Gly Ile Thr Asn Ile Asn Cys Ser Gly 20 25 30

His Ile Trp Val Glu Pro Ala Thr Ile Phe Lys Met Gly Met Asn Ile . 40

Ser Ile Tyr Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu 55

His Phe Tyr Lys Asn Gly Ile Lys Glu Arg Phe Gln Ile Thr Arg Ile 75 70

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Ala Ser Met Tyr Cys Thr Ala Glu Cys Pro Lys His Phe Gln Glu Thr 105

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Asp	Glu 130	Val	Thr	Cys	Val	Ile 135	Tyr	Glu	Tyr	Ser	Gly 140	Asn	Met	Thr	Cys
Thr 145	Trp	Asn	Ala	Gly	Lys 150	Leu	Thr	Tyr	Ile	Asp 155	Thr	Lys	Tyr	Val	Val 160
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Gln	Leu 210	Gln	Ile	His	Leu	Asp 215	Asp	Ile	Val	Ile	Pro 220	Ser	Ala	Ala	Val
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Glu Leu Met Asn Asn Asn Ser Ser Glu Gln Val Leu Tyr Val Asp Pro 355 360 365

Met Ile Thr Glu Ile Lys Glu Ile Phe Ile Pro Glu His Lys Pro Thr 370 375 380

Asp Tyr Lys Lys Glu Asn Thr Gly Pro Leu Glu Thr Arg Asp Tyr Pro 385 390 395 400

Gln Asn Ser Leu Phe Asp Asn Thr Thr Val Val Tyr Ile Pro Asp Leu 405 410 415

Asn Thr Gly Tyr Lys Pro Gln Ile Ser Asn Phe Leu Pro Glu Gly Ser 420 425 430

His Leu Ser Asn Asn Glu Ile Thr Ser Leu Thr Leu Lys Pro Pro 435 440 445

Val Asp Ser Leu Asp Ser Gly Asn Asn Pro Arg Leu Gln Lys His Pro 450 455 460

Asn Phe Ala Phe Ser Val Ser Ser Val Asn Ser Leu Ser Asn Thr Ile 465 470 . 475 480

Phe Leu Gly Glu Leu Ser Leu Ile Leu Asn Gln Gly Glu Cys Ser Ser 485 490 495

Pro Asp Ile Gln Asn Ser Val Glu Glu Glu Thr Thr Met Leu Glu
500 505 510

Asn Asp Ser Pro Ser Glu Thr Ile Pro Glu Gln Thr Leu Leu Pro Asp 515 520 525

Glu Phe Val Ser Cys Leu Gly Ile Val Asn Glu Glu Leu Pro Ser Ile 530 535 540

Asn Thr Tyr Phe Pro Gln Asn Ile Leu Glu Ser His Phe Asn Arg Ile 545 550 555 560

Ser Leu Leu Glu Lys 565

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Met Asn Gln Val Thr Ile Gln Trp Asp Ala Val Ile Ala Leu Tyr Ile

Leu Phe Ser Trp Cys His Gly Gly Ile Thr Asn Ile Asn Cys Ser Gly 20 25

His Ile Trp Val Glu Pro Ala Thr Ile Phe Lys Met Gly Met Asn Ile 35

Ser Ile Tyr Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu 55

<sup>&</sup>lt;210> 25 <211> 356 <212> PRT <213> Homo sapiens

<sup>&</sup>lt;400> 25

His Phe Tyr Lys Asn Gly Ile Lys Glu Arg Phe Gln Ile Thr Arg Ile Asn Lys Thr Thr Ala Arg Leu Trp Tyr Lys Asn Phe Leu Glu Pro His Ala Ser Met Tyr Cys Thr Ala Glu Cys Pro Lys His Phe Gln Glu Thr Leu Ile Cys Gly Lys Asp Ile Ser Ser Gly Tyr Pro Pro Asp Ile Pro Asp Glu Val Thr Cys Val Ile Tyr Glu Tyr Ser Gly Asn Met Thr Cys Thr Trp Asn Ala Gly Lys Leu Thr Tyr Ile Asp Thr Lys Tyr Val Val His Val Lys Ser Leu Glu Thr Glu Glu Glu Gln Gln Tyr Leu Thr Ser Ser Tyr Ile Asn Ile Ser Thr Asp Ser Leu Gln Gly Gly Lys Lys Tyr Leu Val Trp Val Gln Ala Ala Asn Ala Leu Gly Met Glu Glu Ser Lys Gln Leu Gln Ile His Leu Asp Asp Ile Val Ile Pro Ser Ala Ala Val Ile Ser Arg Ala Glu Thr Ile Asn Ala Thr Val Pro Lys Thr Ile Ile Tyr Trp Asp Ser Gln Thr Thr Ile Glu Lys Val Ser Cys Glu Met Arg Tyr Lys Ala Thr Thr Asn Gln Thr Trp Asn Val Lys Glu Phe Asp Thr Asn Phe Thr Tyr Val Gln Gln Ser Glu Phe Tyr Leu Glu Pro Asn Ile Lys Tyr Val Phe Gln Val Arg Cys Gln Glu Thr Gly Lys Arg Tyr Trp Gln Pro Trp Ser Ser Leu Phe Phe His Lys Thr Pro Glu Thr Val Pro

305 310 315 320

Gln Val Thr Ser Lys Ala Phe Gln His Asp Thr Trp Asn Ser Gly Leu 325 330 335

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Gly Ser Tyr Cys 355

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<211> 384

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Leu Phe Ser Trp Cys His Gly Gly Ile Thr Asn Ile Asn Cys Ser Gly 20 25 30

His Ile Trp Val Glu Pro Ala Thr Ile Phe Lys Met Gly Met Asn Ile 35 40 45

Ser Ile Tyr Cys Gln Ala Ala Ile Lys Asn Cys Gln Pro Arg Lys Leu 50 55 60

His Phe Tyr Lys Asn Gly Ile Lys Glu Arg Phe Gln Ile Thr Arg Ile 65 70 75 80

Asn Lys Thr Thr Ala Arg Leu Trp Tyr Lys Asn Phe Leu Glu Pro His 85 90 95

Ala Ser Met Tyr Cys Thr Ala Glu Cys Pro Lys His Phe Gln Glu Thr 100 105 110

Leu Ile Cys Gly Lys Asp Ile Ser Ser Gly Tyr Pro Pro Asp Ile Pro 115 120 125

Asp Glu Val Thr Cys Val Ile Tyr Glu Tyr Ser Gly Asn Met Thr Cys 130 135 140

Thr Trp Asn Ala Gly Lys Leu Thr Tyr Ile Asp Thr Lys Tyr Val Val 145 150 155 160

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Gln	Leu 210	Gln	Ile	His	Leu	Asp 215	Asp	Ile	Val	Ile	Pro 220	Ser	Ala	Ala	Val
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His Val Lys Ser Leu Glu Thr Glu Glu Glu Gln Gln Tyr Leu Thr Ser

Thr Val Ala Ser Ile Ser Thr Gly His Leu Thr Ser Asp Asn Arg Gly 340 345 350

Gln Val Thr Ser Lys Ala Phe Gln His Asp Thr Trp Asn Ser Gly Leu

330

Asp Ile Gly Leu Leu Gly Met Ile Val Phe Ala Val Met Leu Ser 355 360 365

Ile Leu Ser Leu Ile Gly Ile Phe Asn Arg Ser Phe Pro Asn Trp Asp 370 375 380

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<sup>&</sup>lt;212> PRT

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Asp Met Trp Val Glu Pro Gly Glu Ile Phe Gln Met Gly Ile Asn Val 35 40 45

Ser Ile Tyr Cys Gln Glu Ala Leu Lys His Cys Arg Pro Arg Asn Leu 50 55 60

Tyr Phe Tyr Lys Asn Gly Phe Lys Glu Glu Phe Asp Ile Thr Arg Ile 65 70 75 80

Asn Arg Thr Thr Ala Arg Ile Trp Tyr Lys Gly Phe Ser Glu Pro His
85 90 95

Ala Tyr Met His Cys Thr Ala Glu Cys Pro Gly His Phe Gln Glu Thr 100 105 110

Leu Ile Cys Gly Lys Asp Ile Ser Ser Gly His Pro Pro Asp Ala Pro 115 120 125

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Thr Trp Asn Thr Gly Lys Pro Thr Tyr Ile Asp Thr Lys Tyr Ile Val 145 150 155 160

His Val Lys Ser Leu Glu Thr Glu Glu Glu Gln Tyr Leu Ala Ser 165 170 175

Ser Tyr Val Lys Ile Ser Thr Asp Ser Leu Gln Gly Ser Arg Lys Tyr 180 185 190

Leu Val Trp Val Gln Ala Val Asn Ser Leu Gly Met Glu Asn Ser Gln
195 200 205

Gln Leu His Val His Leu Asp Asp Ile Val Ile Pro Ser Ala Ser Ile 210 215 220

Ile Ser Arg Ala Glu Thr Thr Asn Asp Thr Val Pro Lys Thr Ile Val 225 230 235 240

Tyr Trp Lys Ser Lys Thr Met Ile Glu Lys Val Phe Cys Glu Met Arg Tyr Lys Thr Thr Asn Gln Thr Trp Ser Val Lys Glu Phe Asp Ala Asn Phe Thr Tyr Val Gln Gln Ser Glu Phe Tyr Leu Glu Pro Asp Ser Lys Tyr Val Phe Gln Val Arg Cys Gln Glu Thr Gly Lys Arg Asn Trp Gln Pro Trp Ser Ser Pro Phe Val His Gln Thr Ser Gln Glu Thr Gly Lys Arg Asn Trp Gln Pro Trp Ser Ser Pro Phe Val His Gln Thr Ser Gln Thr Val Ser Gln Val Thr Ala Lys Ser Ser His Glu Pro Gln Lys Met Glu Met Leu Ser Ala Thr Ile Phe Arg Gly His Pro Ala Ser Gly Asn His Gln Asp Ile Gly Leu Leu Ser Gly Met Val Phe Leu Ala Ile Met Leu Pro Ile Phe Ser Leu Ile Gly Ile Phe Asn Arg Ser Leu Arg Ile Gly Ile Lys Arg Lys Val Leu Leu Met Ile Pro Lys Trp Leu Tyr Glu Asp Ile Pro Asn Met Glu Asn Ser Asn Val Ala Lys Leu Leu Gln Glu Lys Ser Val Phe Glu Asn Asp Asn Ala Ser Glu Gln Ala Leu Tyr Val Asp Pro Val Leu Thr Glu Ile Ser Glu Ile Ser Pro Leu Glu His Lys Pro Thr Asp Tyr Lys Glu Glu Arg Leu Thr Gly Leu Leu Glu Thr 

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PC17 US 01/31634

A. CLASSIFICATION OF SUBJECT MATTER
I PC 7 C12N15/12 C07K14/705 C07K14/715

According to International Patent Classification (IPC) or to both national classification and IPC

#### **B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols) I PC 7 C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

WPI Data, PAJ, EPO-Internal, MEDLINE, CHEM ABS Data, BIOSIS, SEQUENCE SEARCH, EMB

Category °	Citation of document, with indication, where appropriate, of	the relevant passages	Relevant to claim No.
Υ	WO 98 31811 A (DONALDSON DEBR INST (US); COLLINS MARY (US); 23 July 1998 (1998-07-23) the whole document	A D ;GENETICS NEBEN TA)	1,5-7, 12-35
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χ Furth	er documents are listed in the continuation of box C.	X Patent family members are listed	d in annex.
'A" documer consider filing da filing da filing da filing da citation 'O" documer other m'P" documer	nt which may throw doubts on priority claim(s) or s cited to establish the publication date of another or other special reason (as specified) nt referring to an oral disclosure, use, exhibition or	"T" later document published after the in or priority date and not in conflict wit cited to understand the principle or t invention  "X" document of particular relevance; the cannot be considered novel or canninvolve an inventive step when the d  "Y" document of particular relevance; the cannot be considered to involve an indocument is combined with one or ments, such combination being obvi in the art.  "&" document member of the same paten	h the application but heory underlying the claimed invention of the considered to occument is taken alone claimed invention nventive step when the occupant of the such docuous to a person skilled
	ctual completion of the international search	Date of mailing of the international se	
9	August 2002	1 3. 11. 02	2
lame and m	ailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,	Authorized officer  Armandola, E	

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C.(Continua	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
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L	DATABASE EMBL [Online] Sequence 91 from patent W00193983, 13 February 2002 (2002-02-13) BAKER K.P. ET AL.: "Secreted transmembrane polypeptides and nucleic acids encoding the same." Database accession no. AX358838 XP002209362	
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Int

Box I O	bservations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Interna	ational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X Cl	aims Nos.: 28,30-35 cause they relate to subject matter not required to be searched by this Authority, namely:
h	lthough claims 28 and 30-35 are directed to a method of treatment of the uman/animal body, the search has been carried out and based on the alleged ffects of the compound/composition.
be	aims Nos.: 5,12-15,18, 27-30, 32-35 recause they relate to parts of the International Application that do not comply with the prescribed requirements to such extent that no meaningful International Search can be carried out, specifically:
	ee FURTHER INFORMATION sheet PCT/ISA/210
3. Cla	aims Nos.:
	cause they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II O	bservations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Interna	tional Searching Authority found multiple inventions in this international application, as follows:
S	ee additional sheet
	all required additional search fees were timely paid by the applicant, this International Search Report covers all archable claims.
2. As of	all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment any additional fee.
3. As	only some of the required additional search fees were timely paid by the applicant, this International Search Report vers only those claims for which fees were paid, specifically claims Nos.:
res	required additional search fees were timely paid by the applicant. Consequently, this International Search Report is stricted to the invention first mentioned in the claims; it is covered by claims Nos.:  ,5,12-35(partially), 6,7 (completely)
Remark on	Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1, 5, 12-35 (partially), 6,7 (completely)

An isolated polypeptide HPR1 comprising the amino acid sequence of SEQ. ID. NO: 4, subsequences thereof, fusions of subsequences of SEQ. ID. NO: 4, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequences SEQ. ID. NO: 3, SEQ. ID. NO: 5 and subsequences thereof, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

2. Claims: 1,5,12-35 (all partially)

HPR1 amino acid sequences comprising fragments of SEQ. ID. NO: 4 fused to fragments of SEQ. ID. NO: 1, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

3. Claims: 1,5,12-35 (all partially)

HPR1 amino acid sequences comprising fragments of SEQ. ID. NO: 4 fused to fragments of SEQ. ID. NO: 2, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

4. Claims: 1,5,12-35 (all partially)

HPR1 amino acid sequences comprising fragments of SEQ. ID. NO: 4 fused to fragments of SEQ. ID. NO: 10, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

5. Claims: 1,5,12-35 (all partially)

HPR1 amino acid sequences comprising fragments of SEQ. ID. NO: 4 fused to fragments of SEQ. ID. NO: 11, isolated polynucleotides encoding the polypeptides, expression

vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

6. Claims: 2 and 8 (completely), 5 and 12-35 (partially)

An isolated polypeptide HPR1 comprising the amino acid sequence of SEQ. ID. NO: 12, subsequences thereof, fusions of subsequences of SEQ. ID. NO: 12, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequence SEQ. ID. NO: 28, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

7. Claims: 3, 5,9, 12-35 (all partially)

An isolated polypeptide HPR2 comprising the amino acid sequence SEQ. ID. NO: 23, subsequences thereof, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequence SEQ. ID. NO: 22, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

8. Claims: 3, 5,9, 12-35 (all partially)

An isolated polypeptide HPR2 comprising the amino acid sequence SEQ. ID. NO: 25, subsequences thereof, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequence SEQ. ID. NO: 24, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

9. Claims: 3, 5,12-35 (all partially)

HPR2 amino acid sequences comprising fragments of SEQ. ID. NO: 23 fused to SEQ. ID. NO: 16 or to fragments of SEQ. ID. NO: 16 or to SEQ. ID. NO: 17 or to SEQ. ID. NO: 18, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

10. Claims: 4, 11 (completely), 5 and 12-35 (partially)

An isolated polypeptide HPR2 comprising the amino acid sequence SEQ. ID. NO: 27, subsequences thereof, isolated polynucleotides encoding the polypeptides or comprising the nucleotide sequence SEQ. ID. NO: 29, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

11. Claims: 5, 9 and 12-35 (all partially)

Polynucleotide sequences comprising SEQ. ID. NO: 19 or SEQ. ID. NO: 20, isolated polynucleotides encoding the polypeptides, expression vectors and host cells containing these sequences, processes for producing the polypeptides, antibodies binding to the polypepties, methods for identifying modulators, methods of treatment utilizing the polypeptides.

Continuation of Box I.2

Claims Nos.: 5,12-15,18, 27-30, 32-35

Present claims 5,12-15,18, 27-30, 32-35 relate to an extremely large number of possible compounds/products. Support within the meaning of Article 6 PCT and/or disclosure within the meaning of Article 5 PCT is to be found, however, for only a very small proportion of the compounds/products. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible. Consequently, the search has been carried out for those parts of the claims which appear to be supported and disclosed, namely those parts relating to the DNAs or proteins for which a sequence listing has been provided and to antibodies recognizing the claimed proteins.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

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